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(57) Abstract

cDNAs coding for an infectious Western Equine Encephalitis virus (WEE) and infectious Venezuelan Equine Encephalitis virus variant IE (VEE IE) are disclosed in addition to cDNA coding for the structural proteins of Venezuelan Equine Encephalitis virus variant IIIA (VEE IIIA). Novel attenuating mutations of WEE and VEE IE and their uses are described. Also disclosed are attenuated chimearic alphaviruses and their uses.

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TITLE OF THE INVENTION

LIVE ATTENUATED VIRUS VACCINES FOR EQUINE ENCEPHALITIS VIRUSES

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INTRODUCTION

Western equine encephalitis (WEE), eastern equine encephalitis (EEE) and Venezuelan equine encephalitis virus (VEE) are members of the alphavirus genus of the family Togaviridae which is comprised of a large group of mosquito-borne RNA viruses found throughout much of the world. The viruses normally circulate among 20 rodent or avian hosts through the feeding activities of a variety of mosquitoes. Epizootics occur largely as a result of increased mosquito activity after periods of increased rainfall. Western equine encephalitis virus (WEE) was first recognized in 1930 and causes periodic outbreaks of disease in equines. 25 The virus has been detected over much of the western hemisphere from Argentina north to the more temperate regions of central Canada (For a review, see Reisen and Monath [1988] in The Arboviruses: Epidemiology and Ecology, Vol. V. CRC Press, Inc. Boca Raton). Similarly, EEE was first isolated in Virginia and New Jersey in 1933 (Ten Broeck, C. et al. [1935] J. Exp. Med. 62:677) and is now known to be focally endemic throughout much of the northern portion of South

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America, Central America and the eastern part of Mexico and the United States. Venezuelan equine encephalitis virus has six serological subtypes (I-Two of these subtypes, I and III have multiple variants, two of these variants are of particular interest in this application, variant IE, and variant IIIA also called Mucambo virus. A live, attenuated vaccine (TC-83) for VEE IA/B has been used for immunization equines and at-risk laboratory and field personnel (Birge et al. [1961] Am. J. Hyg. 73:209-218; Pittman et al. [1996] Vaccine 14:337-343). The vaccine was credited with helping to limit the northward spread of a serious epizootic of VEE originating in South America in the late 1960's. 15 However, the VEE I/AB vaccines have not yet been licensed by the Food and Drug Administration and have been shown to be effective in preventing disease from VEE IA/B infection only. The current VEE vaccines do not adequately protect against the VEE IE variant or the VEE IIIA variant, as disease has occurred in 20 laboratory workers successfully vaccinated with a vaccine derived from VEE IA/B. In addition, recent unprecedented outbreaks of VEE IE in populations of horses in Mexico indicate a need for a VEE IE vaccine. 25 The lack of adequate cross protection with existing IA/B vaccines documents the need for a VEE IE-specific

The vaccines currently in veterinary use for WEE, EEE and VEE IA/B throughout the United States and 30 Canada are formalin-inactivated preparations.

Inactivated vaccines for EEE and WEE are also available for use by at-risk laboratory personnel.

These inactivated vaccines are poorly immunogenic, require multiple inoculations with frequent boosters and generally result in immunity of short duration.

and a VEE IIIA-specific vaccine.

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The shortcomings of the available vaccines indicate a need for the development of new vaccines of high immunogenicity which induce a longer lasting immunity for protection against WEE, EEE and VEE subtypes IE and IIIA.

SUMMARY OF THE INVENTION

The present invention satisfies the need mentioned above.

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In this application are described live attenuated vaccines for WEE, EEE, VEE IE and VEE IIIA which may provide higher level immunity in humans and equines for many years, and possibly for life. In addition, very large numbers of vaccine doses can be produced from significantly less starting materials than is 15 possible with the existing inactivated products. The vaccine preparations of the present invention comprise full-length cDNA copies of the genomes of WEE or VEE IE which have been altered such that the RNA produced from the cDNA, and the virus produced therefrom is attenuated and useful as a live vaccine for human and veterinary use. The vaccine preparations for VEE IIIA and EEE are novel chimeric viruses which include the newly discovered structural protein genes of VEE IIIA.

The classic methods of deriving live-attenuated vaccines (blind passage in cell cultures) generally result in heterogeneous and undefined products, hence recent attempts to make live vaccines for alphaviruses have relied on genetic engineering procedures.

The alphavirus genome is a single-stranded, positive-stranded RNA approximately 11,400 nucleotides in length. The 5' two-thirds of the genome consist of a non-coding region of approximately 48 nucleotides followed by a single open reading frame of approximately 7,500 nucleotides which encodes the

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viral replicase/transcriptase. The 3' one-third of the genome encodes the viral structural proteins in the order C-E3-E2-6K-E1, each of which are derived by proteolytic cleavage of the product of a single open reading frame of approximately 3700 nucleotides. sequences encoding the structural proteins are transcribed as a 26S mRNA from an internal promoter on the negative sense complement of the viral genome. The nucleocapsid (C) protein possesses autoproteolytic activity which cleaves the C protein from the 10 precursor protein soon after the ribosome transits the junction between the C and E3 protein coding sequence. Subsequently, the envelope glycoproteins E2 and E1 are derived by proteolytic cleavage in association with 15 intracellular membranes and form heterodimers. initially appears in the infected cell as a precursor, pE2, which consists of E3 and E2. After extensive glycosylation and transit through the endoplasmic reticulum and the golgi apparatus, E3 is 20 cleaved from E2 by furin-like protease activity at a cleavage site having a consensus sequence of RX(K/R)R, with X being one of many amino acids present in the different viruses, and with the cleavage occuring after the last arginine residue. Subsequently, the 25 E2/E1 complex is transported to the cell surface where it is incorporated into virus budding from the plasma membrane (Strauss and Strauss [1994] Microbiological Rev. 58: 491-562). All documents cited herein supra and infra are hereby incorporated in their entirety by 30 reference thereto.

Because the genome of alphavirus is a positivestranded RNA, and infectious upon transfection of cells in culture, an "infectious clone" approach to vaccine development is particularly suitable for the alphaviruses. In this approach, a full-length cDNA clone of the viral genome is constructed downstream from a RNA polymerase promoter, such that RNA which is equivalent to the viral genome can be transcribed from the DNA clone *in vitro*. This allows site-directed mutagenesis procedures to be used to insert specific mutations into the DNA clone, which are then reflected in the virus which is recovered by transfection of the RNA.

Previous work with infectious clones of other 10 alphaviruses has demonstrated that disruption of the furin cleavage site results in a virus which incorporates pE2 into the mature virus. Davis et al. (1995, supra) found that disruption of the furin cleavage site in an infectious clone of VEE is a lethal mutation. Transfection of BHK cells with RNA 15 transcribed from this mutant clone resulted in the release of non-infectious particles. However, a low level of infectious virus was produced which contained secondary suppressor mutations such that virus 20 containing pE2 was fully replication competent and subsequently shown to be avirulent but capable of elliciting immunity to lethal virus challenge in a variety of animal species.

The genetic basis for attenuation of the VEE TC83 vaccine and certain laboratory strains of VEE virus
have been studied extensively and has led to the
development of improved live, attenuated vaccine
candidates (Kinney et al. 1993, supra, Davis et al.
1995, supra). The approach used in this application
is similar to that used for VEE, however, following
the VEE example exactly did not result in an adequate
vaccine for WEE. Changes in the procedure used for
VEE were required, none of which could have been
predicted from the VEE work, in order to produce the
attenuated live WEE virus of the present invention.

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Based upon a comparison of the structural protein gene sequences of WEE and other alphaviruses, the probable furin cleavage site of WEE strain CBA/87 virus is RRPKR. The presence of the extra arginine when compared to the conscensus (RX(R/K)R) alphavirus cleavage site indicated that the cleavage at this site might be more complex than that observed with VEE virus. It was necessary therefore to prepare two deletion mutations in the E3-E2 cleavage site of the full-length clone, one which lacks five amino acids and one which lacks four amino acids since it was unknown which mutation, if any, would produce an attenuated virus. The residual arginine in the fulllength clone lacking only four amino acids was of concern due to the possibility that other mutations might arise due to the presence of the extra arginine resulting in cleavage by cellular proteases at that site and producing an apparently wild type virus with respect to cleavage of pE2.

Transfection of cultured cells with RNA transcribed from an infectious clone of WEE lacking the furin cleavage site yielded viruses which contained the pE2 of WEE in the mature virus but which were not replication competent. During intracellular replication of the RNA, mutations arise at low frequency, resulting in a small number of replication competent virus. Sequence analysis of these viruses has shown that the lethal effect of the deletion mutations was alleviated by the appearance of second site mutations in the E2 glycoprotein. These viruses are attenuated in mice when administered by subcutaneous or intracranial inoculation. The mice produce high titer neutralizing and ELISA antibody and are protected against a lethal challenge of parental virulent WEE virus.

Therefore, in one aspect of the invention, the invention pertains to the isolation of a cDNA sequence coding for an infectious western equine encephalitis (WEE) virus RNA transcript. DNA representing the entire genome, not previously available, was prepared by polymerase chain reaction using a series of primer pairs based upon the partial genome sequences previously deposited in Genbank. The 5' and 3' ends of the viral genome were unknown and difficult to obtain. The terminal sequence was necessary for efficient replication of the virus since substitution of ends from a similar virus with similar but not identical sequences resulted in an extremely attenuated virus. In order to determine the correct 15 sequence at the 5' end, a protocol called rapid amplification of cDNA ends (5'-RACE) was used. full length infectious clone is useful in the production of virulent WEE virus, and introducing and testing attenuating mutations. The production of virulent virus is essential for a formal measure of 20 the degree of attenuation achieved with candidate attenuating mutations and a formal determination of the rate at which reversion to virulence might occur.

In another aspect of the invention, the invention

25 pertains to the isolation of a cDNA sequence coding
for an infectious Venezuelan equine encephalitis virus
IE variant (VEE IE) virus RNA transcript (SEQ ID NO:
2). Using oligonucleotides specific to genomic RNA of
a VEE IE isolate (GenBank accession no. U34999)

30 (Oberste, et al. [1996] Virology 219:314-320),
reverse transcriptase polymerase chain reaction was
carried out to generate numerous cDNA fragments which
were subsequently cloned and used to assemble fulllength cDNA of VEE IE. The full length infectious

35 clone is useful, for example, in the production of

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virulent VEE IE virus, and introducing and testing attenuating mutations.

In the case of VEE IIIA, the structural protein genes were removed from a full length clone of VEE IA/B and replaced by the VEE IIIA structural genes. The IIIA structural gene sequences were prepared by RT-PCR and include the IIIA 26S promoter and the 3' nontranslated region (3'NTR) flanking the sequences for the structural proteins (SEQ ID NO:3). The VEE 10 IIIA 3' NTR was then replaced with the VEE IA/B NTR, and the modified sequence was cloned back into the VEE IA/B full length clone. The result was a clone in which the nonstructural protein gene sequences were from VEE IA/B and the structural protein gene 15 sequences from VEE IIIA. The virus produced from this chimeric clone replicated efficiently in cell culture, and proved to be completely attenuated in mice. In addition, it was highly immunogenic and protected the vaccinated mice against challenge with virulent, wild-20 type Mucambo virus (VEE IIIA).

Portions of the cDNA sequences described above are useful as probes to diagnose the presence of virus in samples, and to define naturally occuring variants of the virus. These cDNAs also make available polypeptide sequences of WEE antigens, EEE antigens, VEE IE antigens, and VEE IIIA structural polypeptide antigens encoded within the respective genomes and permits the production of polypeptides which are useful as standards or reagents in diagnostic tests and/or as components of vaccines. Antibodies, both polyclonal and monoclonal, directed against WEE epitopes, EEE epitopes, VEE IE epitopes, or VEE IIIA epitopes contained within these polypeptide sequences are also useful for diagnostic tests, as therapeutic agents, and for screening of antiviral agents.

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Accordingly, with respect to polynucleotides, some aspects of the invention are: a purified WEE polynucleotide; a recombinant WEE polynucleotide; a recombinant polynucleotide comprising a sequence derived from a WEE genome or from WEE cDNA; a recombinant polynucleotide encoding an epitope of WEE; a recombinant vector containing any of the above recombinant polynucleotides, and a host cell transfected with any of these vectors.

Other aspects of the invention are: a purified VEE IE polynucleotide; a recombinant VEE IE polynucleotide; a recombinant polynucleotide comprising a sequence derived from a VEE IE genome or from VEE IE cDNA; a recombinant polynucleotide encoding an epitope of VEE IE; a recombinant vector containing any of the above recombinant polynucleotides, and a host cell transformed with any of these vectors.

In a further aspect of the invention is a complete sequence of the VEE subtype IIIA structural protein genes useful for diagnostics and vaccine development. Also provided is a chimeric virus containing the structural sequences of VEE IIIA, which is completely attenuated and provides protection against challenge with VEE IIIA virulent virus for use as a vaccine

Another aspect of the invention is a single-stranded DNA sequence comprising a cDNA clone coding for an infectious WEE, the cDNA clone including at least one attenuating mutation therein, the RNA produced from transcription of the cDNA and the virus particles produced from the RNA in a host cell for use as a vaccine.

In another aspect of the invention there is provided a full length WEE cDNA clone containing a

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defined deletion mutation useful for attenuating the virus for the identification of suppressor mutations in the virus. The attenuated virus with the cleavage deletion and suppressor mutations are useful as a means to generate an attenuated, live WEE virus vaccine for veterinary and human use.

In a further aspect of the invention is provided a chimeric virus containing nonstructural protein gene sequences from WEE and structural protein gene

10 sequences from any alphaviruses including but not limited to Aura, Barmah Forest, Bebaru, Cabassou, Chikungunya, eastern equine encephalitis, Everglades, Fort Morgan, Getah, Highlands J, Kyzylagach, Mayaro, Middelburg, Mucambo, Ndumu, O'nyong-nyong, Pixuna,

15 Ross River, Sagiyama, Semliki Forest, SAAR87, Sindbis, Tonate, Una, Venezuelan equine encephalitis, Whataroa, which could be used as a means for attenuating virulent alphaviruses, and vaccine production against other alphaviruses.

Taking advantage of the close evolutionary relationship between WEE and eastern equine encephalitis virus (EEE), a chimeric virus has been constructed in which the structural protein genes of EEE have been inserted into the infectious clone in place of the WEE structural protein genes. The resulting virus is fully replication competent, attenuated, and elicits an immune response in mice which is protective against a lethal challenge with virulent EEE virus.

In addition, depending on the non-WEE sequences substituted for the WEE structural genes, another aspect of the invention includes a means for expressing antigens of other alphaviruses as chimeric alphaviruses for use as potential vaccines for human and veterinary use.

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In another aspect of the invention there is provided a full length infectious VEE IE cDNA clone containing a cleavage deletion useful in the identification of suppressor mutations in the virus, the RNA produced from the cDNA and the virus produced from the RNA. The virus with the cleavage deletion and suppressor mutations is useful as a means to generate an attenuated, live VEE IE vaccine virus for veterinary and human use.

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In a further aspect of the invention is provided a chimeric virus containing nonstructural sequences from VEE IE and structural sequences from other alphaviruses which could be used as a means of attenuating virulent alphaviruses.

In addition, depending on the non-VEE IE sequences substituted for the structural sequences of VEE IE, another aspect of the invention includes a means to express antigens of other alphaviruses as chimeric alphaviruses as potential vaccines for human and veterinary use.

In a further aspect of the invention, there is provided a vaccine protective against WEE, the vaccine comprising live attenuated WEE virus in an amount effective to elicit protective antibodies in an animal to WEE and a pharmaceutically acceptable diluent, carrier, or excipient.

In yet a further aspect of the invention, there is provided a vaccine protective against VEE IE, the vaccine comprising live attenuated VEE IE virus in an amount effective to elicit protective antibodies in an animal to VEE IE and a pharmaceutically acceptable diluent, carrier, or excipient.

In another aspect of the invention, there is provided a bivalent vaccine protective against WEE and VEE IE, the vaccine comprising both attenuated WEE and

attenuated VEE IE in an amount effective to elicit protective antibodies in an animal to both WEE and VEE IE and a pharmaceutically acceptable diluent. In addition, it is possible that this vaccine will provide short lived protection against other alphaviruses (Cole and McKinney [1971] Inf. Immunity 4:37-43).

In yet another aspect of the invention, there is provided an inactivated vaccine produced from the live attenuated virus described above. The attenuated virus of the present invention whether whole virus or chimearic virus can be used in producing inactivated virus vaccines. By using an attenuated virus strain, there is a much greater margin of safety in the event that the produce is incompletely inactivated. Starting with an attenuated strain is also much safer during the manufacturing phase, and allows production under lower biocontainment levels.

20 BRIEF DESCRIPTION OF THE DRAWINGS

These and other features, aspects, and advantages of the present invention will become better understood with reference to the following description and appended claims, and accompanying drawings where:

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Figure 1A, 1B and 1C. Assembly of the full length cDNA clone of western equine encephalitis virus. Polymerase chain reaction products representing the entire genome of WEE virus were prepared with the primer pairs indicated. Each of the products was cloned into pBluescript KS+. Assembly of the full length clone, pWE2000 in pBluescript was carried out as indicated in the figure and described below. The clones are not drawn to scale. In each of the plasmids, the primers used to generate the PCR

products are indicated as are the restriction endonuclease sites used for the assembly.

Figure 2. Polypeptide profiles of western equine encephalitis viruses. Samples of purified virus were analyzed by electrophoresis on 10% polyacrylamide gels and stained with Comassie Brilliant blue. Molecular weight marker (MW) are shown in the first lane and molecular weights are indicated in daltons X 10⁻³. The virus strain is designated above the appropriate lane. Position of contaminating bovine serum albumin is indicated by arrow.

Figure 3. Polypeptide profiles of western equine encephalitis virus (WEE), eastern equine encephalitis (EEE) and chimeric virus, MWE. Samples of purified virus were analyzed by electrophoresis on 10% polyacrylamide gels and stained with Comassie Brilliant blue. Molecular weight markers (MW) are shown and molecular weights are indicated in daltons X 10⁻³. Position of contaminating bovine serum albumin is indicated by arrow.

Figure 4. Derivation of virulent VEE IE clone.

Constructions of cDNA encoding the entire genome of cloned strain 68U201 are shown. The relevent cloning sites and genetic markers are indicated above each clone. The first full-length clone produced that replicated in vitro after transfection of the RNA transcribed from the T7 promoter, was pIE-1006. Further modifications to this clone were carried out in order to obtain pIE-1009 that had the in vitro and in vivo characteristics of the parental, biological isolate, strain 68U201. With a fully virulent clone, pIE-1009 available, specific attenuating mutations

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were introduced into pIE-1009 to generate the attenuated clone, pIE1100. Drawings are not to scale. Shaded areas represent regions of pIE1006 that were replaced or mutated to generate new clones.

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Figure 5. Construction of IAB-IIIA cDNA chimeric clone pV3A-1000.

DETAILED DESCRIPTION

In one embodiment, the present invention relates to a full length cDNA clone of fully virulent WEE virus specified in SEQ ID NO:1, and a full length cDNA clone of fully virulent VEE IE variant specified in SEO ID NO:2.

WEE, strain CBA/87, isolated from the brains of an infected horse in Argentina in 1987 (Bianchi et al., [1987] Am.J. Trop. Med. Hyg. 49:322-328) was used as a parent strain in the instant invention. Any other strain which consistently kills 100% of 5 week old
C57BL6 mice when inoculated subcutaneously can be used

- such as B11, for example. The cDNA clone can be generated by any of a variety of standard methods known in the art. Preferably, DNA representing the entire genome can be prepared by polymerase chain
- reaction using a series of primer pairs based upon the partial genome sequences previously deposited in GenBank. The 5' terminal sequence of the virus may be determined by 5'-RACE basically as described by Frohman et al.([1988] Proc. Natl. Acad. Sci. U.S.A.
- 30 **85**:8998-9002). Assembly of the full length clone can be in a suitable transcription vector such as, for example, pBluescript KS+, using convenient restriction endonuclease sites or the entire genome could be inserted into any plasmid which contains suitable restriction endonuclease cleavage sites for cloning,

an origin of replication so that the plasmid can be propagated in a bacterial host, and a selectable marker gene to maintain the plasmid in the bacterial cell during growth. The DNA sequence preferably has a complementary DNA sequence bonded thereto so that the double-stranded sequence will serve as an active template for RNA polymerase. Hence, the transcription vector preferably comprises a plasmid. When the DNA sequence comprises a plasmid, it is preferred that a unique restriction site be provided 3' (with respect to the virion RNA sequence) to ("down-stream" from) the cDNA clone. This provides a means for linearizing the DNA sequence to enhance the efficiency of transcription of genome-length RNA in vitro.

15 The complete clone is preferably operatively associated with a promoter region such that the promoter causes the cDNA clone to be transcribed in the presence of an RNA polymerase which binds to the promoter. The promoter is positioned on the 5' end 20 (with respect to the virion RNA sequence), or upstream from, the cDNA clone. An excessive number of nucleotides between the promoter sequence and the cDNA clone will result in the inoperability of the construct. Hence, the number of nucleotides between the promoter sequence and the cDNA clone is preferably 25 not more than eight, more preferably not more than than 5, still more preferably not more than 3, and most preferably not more than 1. Exemplary promoters useful in the present invention include, but are not 30 limited to, T3 promoters, T7 promoters, and SP6 promoters. It is preferable that the 5' end of the in vitro transcript not have any additional nucleotides preceding the first nucleotide of the viral sequence. At the 3' end, additional nucleotides can be tolerated 35 in the in vitro transcript but are probably lost when

the virus replicates. In most instances, the poly-dA tract at the 3' end is required for viability of the virus. Selection of the virulent full length clone can be achieved by comparing the LD_{50} of the virus encoded by the cloned cDNA with the LD_{50} of the parent virus used, in the instant example, WEE CBA/87. The ability to produce virulent virus is important; it allows the introduction and testing of attenuation mutations and the attenuated phenotype against a standard: the virulent parent.

Transfection of cells with the RNA transcript coded by the full length genomic cDNA can be achieved by any suitable means, such as, for example, by treating the cells with DEAE dextran, treating the cells with "LIPOFECTIN", and by electroporation.

Togavirus-permissive cells, alphavirus-permissive cells, and WEE-permissive and VEE IE-permissive cells are cells which, upon transfection with the viral RNA transcript, are capable of producing viral particles.

Togaviruses have a broad host range. Examples of such cells include, but are not limited to, Vero cells, baby hamster kidney cells, chicken embryo fibroblast cells, Chinese hamster ovary cells (CHO), mouse L cells, MRC-5 cells, mosquito cells such as C6-36 cells, to name a few.

In the case of VEE IE, an isolate of VEE IE, strain 68U201, was used. Any isolate known to cause disease in man can be chosen. In addition the ability of a strain to have an easily identifiable phenotype such as, for example, the ability to form large plaques in tissue culture on indicator cell monolayers, is helpful. In the present invention, the full length clone of VEE IE was obtained using oligonucleotide primers specific for the VEE IE strain 68U201 sequence. Reverse transcription-polymerase

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chain reaction (RT-PCR) of strain 68U201 viral RNA was carried out to generate numerous cDNA fragments that were subsequently cloned and used to assemble a full-length cDNA in a plasmid such that the cDNA could be precisely transcribed and viral infectious RNA produced. It is also possible to produce the entire genome by polymerase chain reaction by including the promoter sequence in the 5' end primer thereby producing infectious RNA directly from the PCR fragment. It is also possible to transcribe the viral RNA from a plasmid in the cell by transfection with the appropriate plasmid containing a promoter utilized by cellular RNA polymerases, i.e. the CMV promoter.

To determine virulence of the cloned viral genome, mice can be innoculated subcutaneously with 10⁴ plaque forming units of the cloned virus. The clone is considered virulent if all mice die, and not fully virulent if mice do not all die. The LD₅₀ of parent VEE IE strain 68U201 is to 1-2 pfu, therefore, if inoculation of 10,000-fold did not cause lethal disease in all mice, it was considered attenuated.

In another embodiment of the present invnetion is provided a cDNA sequence of the entire 26S region of VEE subtype IIIA as well as the structural protein genes specified in SEQ ID NO:3.

DNA or polynucleotide sequences to which the invention also relates include sequences of at least about 6 nucleotides, preferably at least about 8 nucleotides, more preferably at least about 10-12 nucleotides, most preferably at least about 15-20 nucleotides corresponding, i.e., homologous to or complementary to, a region of the WEE, VEE IE, or VEE IIIA nucleotide sequence. Preferably, the sequence of the region from which the polynucleotide is derived is homologous to or complementary to a sequence which is

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unique to the virus. Whether or not a sequence is unique to the virus can be determined by techniques known to those of skill in the art. For example, the sequence can be compared to sequences in databanks, e.g., GenBank. Regions from which typical DNA sequences may be derived include but are not limited to, for example, regions encoding specific epitopes, as well as non-translated regions.

The derived polynucleotide is not necessarily 10 physically derived from the nucleotide sequence of the alphaviruses, but may be generated in any manner, including for example, chemical synthesis or DNA replication or reverse transcription or transcription, which are based on the information provided by the sequence of bases in the region(s) from which the polynucleotide is derived. In addition, combinations of regions corresponding to that of the designated sequence may be modified in ways known in the art to be consistent with an intended use. The sequences of the present invention can be used in diagnostic assays such as hybridization assays and polymerase chain reaction assays and for the discovery of other alphavirus sequences.

A polypeptide or amino acid sequence derived from 25 the amino acid sequence of alphavirus, refers to a polypeptide having an amino acid sequence identical to that of a polypeptide encoded in the sequence, or a portion thereof wherein the portion consists of at least 2-5 amino acids, and more preferably at least 8-10 amino acids, and even more preferably at least 11-15 amino acids, or which is immunologically identifiable with as a polypeptide encoded in the sequence.

A recombinant or derived polypeptide is not 35 necessarily translated from a designated nucleic acid

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sequence, it may be generated in any manner, including for example, chemical synthesis, or expression of a recombinant expression system.

Once a complete viral genomic cDNA is cloned, attenuation of the virus is possible. An attenuating mutation refers to a nucleotide mutation or amino acid coded for in view of such a mutation which results in a decreased probability of causing disease in its host (i.e., a loss of virulence), in accordance with standard terminology in the art. The attenuating mutation may be a substitution mutation or an in-frame deletion mutation.

Novel WEE attenuating mutations disclosed herein which may be used to carry out the present invention include deletion of five amino acids at the furin 15 cleavage site in combination with a substitution of lysine for glutamic acid at codon 182 at E2, or deletion of five amino acids at the furin cleavage site in combination with a subsitution of lysine for glutamic acid at codon 181 at E2. Additionally, 20 certain mutations placed in the non-coding region at the 5' end of the genome have been found to be attenuating, specifically, a C to T change at nucleotide 7, an A to G change at nucleotide 13, a T to A change at nucleotide 25 and deletion of an A at nucleotide 22. These novel attenuating mutations may be inserted together in a cDNA clone encoding WEE virus resulting in an attenuated WEE which is reflected by 100% survival of mice inoculated by 30 subcutaneous and intracranial routes. Such an attenuated live virus is immunogenic and protective against a lethal virus challenge.

Novel attenuating mutations can be discovered in the VEE IE by introducing mutations which are not reparable by the viral RNA replication process. A

preferable mutation is the deletion of the four amino acids of the furin-like cleavage site between the E3 and E2 proteins. Transfection of the mutant viral genome into cells can result in the suppression of the lethal effect of the deletion mutation due to the error prone process of alphavirus replication. Once efficiently replicating viral progeny is generated they can be detected by plaque assays and analyzed for the presence of pE2 protein which indicates that the virus contains the deletion mutation. Attenuated but yet immunogenic virus with a cleavage deletion mutation and suppressor mutation(s) could be tested for its ability to protect animals from challenge with virulent VEE IE.

15 Attenuating mutations may be introduced into cDNAs encoding live WEE or VEE IE by any suitable means, such as site-directed mutagenesis (Please see e.g., Maniatis, Fritsch and Sambrook, Molecular Cloning: A Laboratory Manual (1982) or DNA Cloning, Volumes I and II (D. N. Glover ed. 1985) or Current Protocols in Molecular Biology, Ausubel, F. M et al. (Eds.) John Wiley & Sons, Inc., for general cloning methods.).

In another embodiment of the present invention is provided a chimeric virus containing nonstructural sequences from one alphavirus and structural sequences from other alphaviruses which could be used as a means of attenuating virulent alphaviruses. By "Structural sequences" as used herein is meant sequences encoding proteins which are required for encapsidation (e.g., packaging) of the viral genome, and include the capsid protein, El glycoprotein, and E2 glycoprotein. By "nonstructural sequences" is meant nonstructural protein sequences, or sequences which encode viral RNA

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polymerase(s) proteins. Viruses from which structural sequences can be used in the chimeric virus using WEE "nonstructural genes" as the backbone clone can include for example, all strains of WEE, EEE, and Sindbis, Aura, Barmah Forest, Bebaru, Bijou Bridge, Cabassou, Chikungunya, Everglades, Fort Morgan, Getah, Highlands J, Kyzylagach, Mayaro, Middelburg, Mucambo, Ndumu, O'nyong-nyong, Pixuna, Ross River, Sagiyama, Semliki Forest, SAAR87, Tonate, Una, Venezuelan Equine Encephalitis, Whataroa, to name a few. Acceptable 10 structural protein genes would include a nucleocapsid protein capable of both packaging the chimeric viral genome and which can interact with the glycoproteins to initiate particle assembly. Chimearic virus is 15 constructed by excision of the structural protein genes of the backbone virus and replacement with the desired structural protein genes from another virus. This can be accomplished in one of several ways. For example, site-directed mutagenesis can be used to 20 excise the structural protein genes and leave a restriction endonuclease digestion site at the point of deletion. The structural protein genes of another alphavirus would then be cloned into that restriction site. Any virus obtained after transfection of cells with RNA transcribed from that clone would by definition be a chimeric virus.

In the case where the first and second viruses are closely related, another method can be used wherein cloned structural cDNA sequences of a second alphavirus can be digested at restriction enzyme sites which both viruses have in common. The cDNA fragments of the second virus can then be cloned into the homologous sites in the first virus structural protein locus such that the resulting structural protein genes of the chimeric are a composite of both. Other

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methods for producing a chimearic virus are known to people in the art (Kuhn et al. [1996] J. Virology 70:7900-7909).

In another embodiment, the attenuated viruses of the present of invention can be used to prepare 5 replicon expression systems. A replicon expression system consists of three components. The first is a replicon which is equivalent to a full length infectious clone from which all of the viral 10 structural proteins have been deleted. A multiple cloning site can be cloned into the site previously occupied by the structural protein genes. Virtually any heterologous gene may be cloned into this cloning Transcription of RNA from the replicon yields 15 an RNA capable of initiating infection of the cell identically to that seen with the full-length infectious virus clone. However, in lieu of the viral structural proteins, the heterologous antigen is expressed. This system does not yield any progeny 20 virus particles because there are no viral structural proteins available to package the RNA into particles.

Particles which appear structurally identical to virus particles can be produced by supplying structural proteins for packaging of the replicon RNA in trans. This is typically done with two helpers. One helper consists of a full length infectious clone from which the nonstructural protein genes and the glycoprotein genes are deleted. The helper retains only the terminal nucleotide sequences, the promoter for subgenomic mRNA transcription and the sequence for the viral nucleocapsid protein. The second helper is identical to the first except that the nucleocapsid gene is deleted and only the glycoprotein genes are retained. The helper RNA's are transcribed in vitro and co-transfected with replicon RNA. Because the

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replicon RNA retains the sequences for packaging by
the nucleocapsid protein, and because the helpers lack
these sequences, only the replicon RNA is packaged by
the viral structural proteins and released from the

5 cell. The particles can then be inoculated into
animals similar to parent virus. The replicon
particles will initiate only a single round of
replication because the helpers are absent, they
produce no progeny virus particles, and express only
the viral nonstructural proteins and the product of
the heterologous gene cloned in place of the
structural proteins. The heterologous gene product is
then detected by the host immune system and
appropriate immune response is then mounted.

The WEE and VEE IE replicons can be used to express heterologous genes of interest as well as a means for expressing antigens or immunogenic proteins and peptides of interest, in vitro or in vivo. The immunogenic protein or peptide, or "immunogen" may be any immunogen suitable for inducing an immune response protective against a pathogen from which the immunogen is derived, including but not limited to microbial, bacterial, protozoal, parasitic, and viral pathogens. For example, the immunogen can be the expression product of any heterologous gene of interest, including influenza hemagglutinin, lassa fever nucleocapsid and glycoproteins, portions of bacterial toxin genes, HIV glycoprotein, Ebola glycoprotein, to name a few.

In yet another embodiment, the present invention provides inactivated virus vaccines produced from live attenuated virus preparations, either as virus with attenuating mutations as has been described for WEE and VEE IE or chimeric viruses described above for EEE and VEE IIIA. The inactivation of live virus is well

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known in the art and can be performed, for example, by the use of formalin. Inactivated attenuated virus vaccine has a greater safety margin both as a final vaccine in case of incomplete inactivation, and during the manufacturing process allowing production under lower biocontainment levels.

Subjects which may be administered the live attenuated or inactivated attenuated viruses and vaccine formulations disclosed herein include both humans and animals (e.g. horse, donkey, pigs, mice, hamster, monkey, birds).

Vaccine formulations of the present invention comprise an immunogenic amount of a live attenuated virus, or a combination of live attenuated viruses as a multivalent vaccine, as disclosed herein in combination with a pharmaceutically acceptable carrier. An "immunogenic amount" is an amount of the attenuated virus sufficient to evoke an immune response, particularly an immune response to the protein or peptide encoded by the heterologous RNA carried by the virus, in the subject to which the virus is administered. An amount of from about 101 to 10⁵ plaque forming units of the live virus per dose is suitable, depending upon the age and species of the subject being treated. Exemplary pharmaceutically acceptable carriers include, but are not limited to, sterile pyrogen-free water and sterile pyrogen-free physiological saline solution.

Administration of the live attenuated viruses disclosed herein may be carried out by any suitable means, including both parenteral injection (such as intraperitoneal, subcutaneous, or intramuscular injection), by in ovo injection in birds, and by topical application of the virus (typically carried in the pharmaceutical formulation) to an airway surface.

Topical application of the virus to an airway surface can be carried out by intranasal administration (e.g. by use of dropper, swab, or inhaler which deposits a pharmaceutical formulation intranasally). Topical application of the virus to an airway surface can also be carried out by inhalation administration, such as by creating respirable particles of a pharmaceutical formulation (including both solid particles and liquid particles) containing the virus as an aerosol suspension, and then causing the subject to inhale the respirable particles. Methods and apparatus for administering respirable particles of pharmaceutical formulations are well known, and any conventional technique can be employed.

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15 In another embodiment, the present invention relates to antibodies specific for the above-described virus. For instance, an antibody can be raised against any of the viral proteins or against a portion thereof. Persons with ordinary skill in the art using 20 standard methodology can raise monoclonal and polyclonal antibodies to a polypeptide of the present invention. Material and methods for producing antibodies are well known in the art (see for example Goding, in, Monoclonal Antibodies: Principles and 25 Practice, Chapter 4, 1986). The antibodies can be used to monitor the presence or activity of alphaviruses and potentially as a therapeutic agent.

In a further embodiment, the present invention relates to a method of detecting the presence of WEE, EEE, VEE IIIA or VEE IE viral infection or antibodies against these viruses, if present, in a sample. Using standard methodology well known in the art, a diagnostic assay can be constructed by coating on a surface (i.e. a solid support) for example, a

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microtitration plate or a membrane (e.g. nitrocellulose membrane), all or a unique portion of WEE, EEE, VEE IIIA or VEE IE virus described above, and contacting it with the serum of a person suspected of having a viral infection. The presence of a resulting complex formed between the virus and antibodies specific therefor in the serum can be detected by any of the known methods common in the art, such as colorimetry or microscopy. This method of detection can be used, for example, for the diagnosis of WEE, EEE, VEE IIIA and VEE IE viral infections.

In yet another embodiment, the present invention relates to a method of detecting the presence of WEE, 15 EEE, VEE IIIA or VEE IE viruses in a sample. Using standard methodology well known in the art, a diagnostic assay can be constructed by coating on a surface (i.e. a solid support) for example, a microtitration plate or a membrane (e.g. 20 nitrocellulose membrane), antibodies specific for WEE and/or VEE IE, and contacting it with serum or tissue sample of a person suspected of having a WEE or VEE IE viral infection. The presence of a resulting complex formed between virus in the serum and antibodies 25 specific therefor can be detected by any of the known methods common in the art, such as fluorescent antibody spectroscopy or colorimetry. This method of detection can be used, for example, for the diagnosis

In another embodiment, the present invention relates to a diagnostic kit which contains WEE, EEE, VEE IE, or VEE IIIA virus and ancillary reagents that are well known in the art and that are suitable for use in detecting the presence of antibodies to WEE and VEE IE in serum or a tissue sample. Tissue samples

of WEE, EEE, VEE IIIA and VEE IE viral infection.

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contemplated can be obtained from birds, monkey, human, or other mammals.

In yet a further embodiment, the present invention relates to DNA or nucleotide sequences for use in detecting the presence or absence of WEE, EEE, VEE IE or VEE IIIA virus using the reverse transcription-polymerase chain reaction (RT-PCR). DNA sequence of the present invention can be used to design primers which specifically bind to the viral RNA for the purpose of detecting the presence, absence, or quantitating the amount of virus. primers can be any length ranging from 7-40 nucleotides, preferably 10-15 nucleotides, most preferably 18-25 nucleotides. Reagents and controls necessary for PCR reactions are well known in the art. 15 The amplified products can then be analyzed for the presence or absence of viral sequences, for example by gel fractionation, with or without hyridization, by radiochemistry, and immunochemical techniques.

In yet another embodiment, the present invention relates to a diagnostic kit which contains PCR primers specific for WEE, EEE, VEE IE or VEE IIIA and ancillary reagents that are well known in the art and that are suitable for use in detecting the presence or absence of WEE, EEE, VEE IE or VEE IIIA in a sample using PCR. Samples contemplated can be obtained from birds, human, or other mammals.

In another embodiment, the present invention relates to a method of reducing WEE, EEE, VEE IE, or VEE IIIA viral infection symptoms in a patient by administering to said patient an effective amount of anti WEE, anti EEE, anti VEE IE, or anti VEE IIIA antibodies, or protective serum from an immunized animal. When providing a patient with antibodies, the dosage of administered agent will vary depending upon

such factors as the patient's age, weight, height, sex, general medical condition, previous medical history, etc. In general, it is desirable to provide the recipient with a dosage of the above compounds which is in the range of from about 1pg/kg to 10 mg/kg body weight of patient, although a lower or higher dosage may be administered.

In another embodiment, the present invention relates to a method for overcoming vaccine interference in alphavirus-immune inquviduals. 10 Alphavirus interference has been documented in animals and people since the 1960's. This phenomenon occurs when a live-attenuated vaccine is administered to animals or people with existing immunity to heterologous alphaviruses. Pre-existing immunity may 15 be acquired by vaccination or infection. This presents a significant limitation to the usefulness of the current live-attenuated alphavirus vaccines, especially since the cross-reactive immunity does not 20 protect adequately against challenge with virulent heterologous alphaviruses. Formalin-inactivated vaccines are not an acceptable alternative as they have significant limitations with regard to the quality and duration of protective immunity and 25 require multiple inoculations and periodic boosters. The attenuated WEE, EEE, VEE IIIA and VEE IE virus vaccines of the present invention contain mutations in the viral glycoprotein sequences that may alter the sequence, conformation, and/or accessibility of crossreactive epitopes. Alterations in epitopes that 30 prevent binding by cross-reactive antibodies may also bypass interference in alphavirus-immune individuals. Eliminating the problem of interference would permit the WEE and VEE IE attenuated virus vaccines to be used in alphavirus-immune animals or people to induce 35

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protective immunity to western equine encephalitis virus virus and/or to Venezuelan equine encephalitis virus variant IE. Long-lasting protective immunity to both parenteral and aerosol challenge would be expected after vaccination with the live-attenuated vaccines of the present invention, and provide an additional advantage over the use of inactivated vaccines which induce short-lived responses that do not protect against mucosal challenge.

Having now described the invention, the following examples are provided to illustrate the present invention, and should not be construed as limiting thereof. In light of the present disclosure, numerous embodiments within the scope of the claims will be apparent to those of ordinary skill in the art.

Attenuated WEE

The following materials and methods were used in the examples that follow.

Viruses and cells. Western equine encephalitis virus, strain CBA/87 (Bianchi et al. 1988) and eastern equine encephalitis virus, strain Fla91-4679 (Mitchell et al.[1992] Science 257:526-7) were grown in BHK cells in EMEM containing 10% fetal bovine serum, 100 U/ml penicillin G and 100 μ g/ml streptomycin. Primary chicken embryo fibroblasts were grown in EMEM containing 5% fetal bovine serum. Baby hamster kidney cells and Vero cells were grown in EMEM containing 10% fetal bovine serum.

<u>cDNA cloning</u>. Genomic RNA was prepared from purified virus by phenol:chloroform extraction and ethanol precipitation. Initially, cDNA was prepared by the method of Gubler and Hoffman ([1983] *Proc*.

- Natl. Acad. Sci. U.S.A. 85:5997-6001). Subsequently, DNA representing the entire genome was prepared by polymerase chain reaction using a series of primer pairs (Table 1, Figure 1) based upon the partial genome sequences previously deposited in GenBank. Each PCR product was cloned into pCRII (Invitrogen). The 5' terminal sequence of CBA/87 virus was determined by 5'-RACE basically as described by Frohman et al. ([1988] Proc. Natl. Acad. Sci. U.S.A.
- 10 **85**:8998-9002). The oligonucleotide, CBA/87 5', consisted of an *Sst*II site, the promoter for bacteriophage T7 RNA polymerase followed by one G and 14 nucleotides of the 5' terminus of CBA/87 terminal sequence was paired with ns1962 and used to amplify
- the terminal 1.9 kb of the WEE genome. Clone pWE2000 representing the entire genome was assembled in pBluescript KS+ through the use of convenient restriction endonuclease sites.
- Table 1. Oligonucleotide Primers for Preparation of WEE PCR Products
 CBA/87 -T7-Sst2

GTCACCGCGGTAATACGACTCACTATAGATAGGGCATGGTATAGAG (SEQ ID NO:4)

- 25 NS1962 TCACCTTATTCTGGAACACATCAG (SEQ ID NO:5) WEE-7 TCGGAGGAAGGCTGATGAAAC (SEQ ID NO:6) WEE-10 TCGGATCCGATGAGAAAATATACGCTCCC (SEQ ID NO:7) WEE-17 GACTGGATCCGCAAACCAGTCCTGTTCTCAGG (SEQ ID NO:8) 30 WEE-18 GCATGGATCCAGCATGATCGGAAATGTCTTGTC (SEO ID NO: 9) WEE-5 TCGGATCCACCGCCAAAATGTTTCCATAC (SEQ ID NO:10) TCGGGATCCCCGGAACATTTGGC (SEQ ID NO:11) WEE-3 CTGCTTTTCATGCTGCATGCC (SEQ ID NO:12) WEE-2 35 WEE-Not CGATGCGGCCGCTTTTTTTTTTTTTTTTTTTTGAAATTTTAAAAAC (SEQ ID NO:13) WEE-CL2 CAGCGTGAAGTCATCGGTAATGCTGCGTGATGGACATTTCAAG
- (SEQ ID NO:14)
 WEE-CL1 CAGCGTGAAGTCATCGGTAATGCTTGATGGACATTTCAAG
 (SEQ ID NO:15)

For ease of subsequent site-directed mutagenesis of the structural protein genes, two cassettes representing the 5' terminal 7.6 kb, plasmid pWE5'-18, and 3' 4.2 kb, plasmid pWE3'-17, of the genome were prepared. Full length clones were assembled by digestion of the pWE5'-18 with BlnI and NotI and insertion of a 4.1 kb BlnI-NotI fragment prepared from the plasmid pWE3'-17 or its mutagenized derivatives.

A cassette containing the structural genes of eastern equine encephalitis virus strain Fla91-4679 was prepared by RT-PCR. The cassette was digested with *BlnI* and *NotI* and the 4.0 kb fragment was ligated to pWE5'-18 which had been similarly digested. The resulting plasmid was designated pMWE-7.

Mutagenesis of the furin cleavage site. 15 Two oligonucleotides, WEE-CL2 and WEE-CL which bracket the presumed furin cleavage site, RRPKR, between the E3 and E2 glycporoteins were used to delete the 5 and 4 codons, respectively. Plasmid pWE3'-17 was used as template for mutagenesis. WEECL2 and WEECL were 20 paired with primer WEE-5 to generate PCR products of 1.4 kb. The PCR products were purified and paired with WEE-3 for 10 cycles of PCR utilizing WE3'-17 as template. Additional WEE-5 primer was added to 500 nM and PCR was carried out for an additional 20 cycles. 25 The 2.3 kb products were purified, digested with BstEII and NcoI and ligated into plasmid pWE3'-17 which had been digested with BstEII and NcoI. Clones containing the mutation were identified by loss of the 30 NgoM1 site in the sequences encoding the furin cleavage site. The sequences of the mutations were confirmed by sequencing and the mutagenized pWE3'-17 cassettes were ligated to pWE5'-18 as described above to yield full length clones pWE2200 and pWE2100, 35 respectively.

Identification of secondary mutations in virus derived from pWE2100 and pWE2200. Virus released from cells electroporated with RNA transcribed from pWE2200 were plaque-purified and grown into stock preparations. Supernatant from cells transfected with RNA transcribed from pWE2100 was diluted 10-fold and inoculated onto BHK cells. The supernatant was collected 48 hours later and the RNA was extracted directly from the culture fluid. For each mutant virus and the parent CBA/87 virus, viral RNA was 10 extracted with Trizol LS (Life Technologies, Inc., Gaithersburg, MD) and the glycoprotein genes of each were amplified by reverse transcription-polymerase chain reaction amplification. The PCR products were 15 purified (PCR Prep, Promega Inc., Madison, WI) and sequenced on an ABI 373 sequencer using fluorescenttagged terminators.

Transcription and transfection. Purified plasmid was digested with NotI, phenol extracted and ethanol 20 precipitated. Typically 0.5-1 ug of linearized DNA was transcribed in vitro by T7 RNA polymerase (Ribomax, Promega, Madison, WI) in the presence of 3 mM m7GpppGp (Pharmacia, Piscataway, NJ). Electroporation of BHK or CEF cells with 0.4 ug of RNA 25 was done as described (Liljestrom et al. [1991] Bio/Technology 9:1356-1361). The cells were then seeded into T-75 flasks with 20 ml of medium and observed for cytopathology at 24 and 36 hours after electroporation. Virus was harvested when the cells displayed significant cytopathology and approximately 50% were detached from the plastic. Virus titers were determined by plaque assay on Vero and BHK cells.

EXAMPLE 1

Preparation of an infectious clone of WEE strain CBA/87.

One goal of this study was to prepare a full 5 length cDNA clone of fully virulent WEE virus such that mutations leading to an attenuated phenotype could be identified. WEE, strain CBA/87, isolated from the brain of an infected horse in Argentina in 1987 (Bianchi et al. [1988] Am. J. Trop. Med. Hyg. 49:322-328), was chosen as the parent virus as it 10 consistently kills 100% of 5 week old C57BL6 mice when inoculated subcutaneously, allowing development of a convenient animal model to assess the relative effects on virulence of the attenuating mutations. Portions of the sequence of several strains of WEE virus had 15 been determined previously and were used as a basis for primers to prepare amplification products representing the entire genome of the CBA/87 which were cloned in pCRII. Full length clones were 20 assembled in pBluescript KS+ using convenient restriction sites as shown in Figure 1. The first infectious clone pWE1000, contained the 5' terminal 20 nucleotides from eastern equine encephalitis virus. This clone produced viable virus which was highly attenuated exhibiting an subcutaneous LD_{50} in mice of 25 approximately 1.2 X 106 PFU compared to the CBA/87 parent virus where LD₅₀ was approximately 22.

A second clone with an authentic WEE 5'-terminus, pWE2000, was used for all subsequent experiments. Transfection of CEF or BHK cells with RNA transcribed from pWE2000 resulted in complete destruction of the monolayers within 36 hours and titers >108 PFU/ml were obtained by infection of either cell type with the resulting virus. Subcutaneous inoculation of 5 week

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old C57/B16 mice with WE2000 results in death within 9 days and the LD_{50} of 75 PFU is only slightly higher than the LD_{50} of 22 PFU of CBA/87 parent virus, Table 2. It should be noted that animals surviving the lower doses of virus challenge gave no serological evidence of infection. This level of virulence for WE2000 virus was considered sufficient to allow for further characterization of the mutations necessary for attenuation of the virus.

Table 2. C57 Black/6 Mice Inoculated Subcutaneously with CBA/87 Parental virus and WE2000 Recombinant Virus

Virus Strain Dose	Mortality	Mean Day to Death (Days)	Prechallenge ELISA	Challenge (S/T)
CBA/87 3X10 ⁵ 3X10 ⁴ 3X10 ³ 3X10 ² 3X10 ¹	10/10 10/10 10/10 7/10 3/10 0/10	8.4 8.9 9.2 9.9 10	- - <100 <100 <100	- - 0/3 0/7 0/10
WE2000 1X10 ⁷ 1X10 ⁶ 1X10 ⁵ 1X10 ⁴ 1X10 ³ 1X10 ² 1X10 ¹	10/10 10/10 10/10 10/10 9/10 3/10 0/10	8.7 9.0 9.2 9.3 10.0	- - - <100 <100 <100 <100	- - - 0/1 0/7 0/10 0/10

EXAMPLE 2

Preparation of cleavage mutants of CBA/87.

Davis et al.([1995] Virology 212:102-110) have demonstrated that deletion of the furin cleavage site between E3 and E2 glycoproteins of VEE virus is a lethal mutation. However, prolonged incubation of cells which had been transfected with RNA derived from

full length clones with the deletion, resulted in the eventual appearance of virus which was replication-competent and attenuated in mice (Davis et al., 1995, supra). Based upon a comparison of the predicted structural protein sequences of WEE and VEE, the probable cleavage site of CBA/87 virus is RRPKR. The presence of the extra arginine when compared to the conscensus (RX(R/K)R) alphavirus cleavage site indicated that the cleavage site of WEE virus might be more complex than that observed with VEE virus. We therefore, prepared two different deletion mutations in the E3-E2 cleavage site of the pWE2000 clone, pWE2100 which lacks five amino acids, RRPKR, and pWE2200 which lacks four amino acids, RPKR.

15 Development of CPE in cells after electroporation of RNA transcribed from pWE2100 and pWE2200 was delayed for 48 to 72 hours compared with pWE2000. In two instances, pWE2100 did not induce significant CPE in the transfected cells despite the fact that 20 approximately 10° PFU/ml were released into the medium of transfected cells. When assayed by plaque formation on Vero cells, both supernatants yielded extremely small plaques after 72 hours which never increased beyond 2 mm in diameter. In contrast. 25 pWE2000 virus yields large plaques after 48 hours which enlarge to approximately 1 cm after 5 days under a 0.5% agarose overlay. The small plaque phenotype of the mutant viruses is stable after 3 passages in Vero or BHK cells, which is the limit to the passage of the 30 virus used in these experiments.

Analysis of the structural proteins of the WE2100 and WE2200 viruses by SDS-PAGE (Figure 2) shows that in each instance, the deletions at the cleavage site result in a virus which lacks E2 protein and contains a larger protein, presumably pE2, indicating that

deletion of the presumed cleavage site eliminated cleavage at this site.

The lack of rapid cytopathology after transfection of BHK cells with transcripts of pWE2200 and pWE2100 suggested that the mutants were non-viable and that the infectious virus subsequently detected by plaque assay was due to secondary mutations arising during the replication of the RNA as reported previously for deletion mutants of VEE (Davis et al., 1995, supra).

Three plaque isolates from WE2200 virus were chosen for further characterization. All isolates grew to high titer and exhibited a small plaque phenotype. The isolates were sequenced over the entire glycoprotein reading frame. As shown in Table 3, isolates 2215 and 2220 have a mutation of Glu to Lys at position 181 of the E2 glycoprotein. Strain WE2219 carries a single Glu to Lys at position 182 of the E2 protein. Strain WE2215 also has a conservative Val to Ala change at position 211 of the E2 glycoprotein. WE2220 has a Glu to Gly change at position 2 of the E1 glycoprotein and a Phe to Ser change at position 382 of the E1 glycoprotein.

25 <u>Table 3. Genotypes of Recombinant WEE Virus Strains</u>

	<u>Virus¹</u>	<u>Cleavage</u> <u>Site</u>	<u>E2°</u>	<u>E1²</u>
30	CBA/87	RRPKR	P(102),E(181),E(182)	E(2),F(257),P(382)
	vWE2100 pWE2102		K(182) K(182)	
35	vWE2215 vWE2219 vWE2220	R R R	K(181), A(211) K(182) K(181)	G(2), S(382)

- 1. p indicates viruses prepared by mutagenesis of infectious clones.
- 2. Amino acid at position indicated in parenthesis.

In order to determine a consensus of the mutations appearing in virus produced from the pWE2100 RNA, the sequences of the glycoprotein genes were determined directly from cDNA prepared by RT-PCR of the genomic RNA extracted from virus released from BHK cells infected with virus released from the transfected cells. Sequence analysis of glycoprotein genes of WE2100 virus from the transfection supernatant revealed only two mutations. As seen previously in WE2219, WE2100 also had a Glu to Lys change at position 182 of the E2 glycoprotein (Table 3).

In order to determine which of the mutations identified in virus released from cells transfected with RNA transcripts from pWE2200 and pWE2100 served 20 as the suppressor of the lethal effect of the cleavage deletion mutation, the mutations were individually placed into the pWE2200 and pWE2100 clones by sitedirected mutagenesis. As shown in Table 4, three subclones of WE2200 were produced, and based upon the 25 ability to induce CPE in BHK cells, it was demonstrated that the Glu to Lys change at position 181 of the E2 glycoprotein was necessary and sufficient to restore the ability of the WE2200 clone to encode replication competent virus. Similarly, placement of the Glu to Lys change at position 182 of the E2 glycoprotein was also sufficient to restore the ability of the WE2100 clone to encode replication competent virus. When the Glu to Lys changes at E2 position 181 or 182 were inserted into the parental 35 infectious clone pWE2000, the resulting virus

exhibited a small plaque phenotype on Vero cells as noted for each of the cleavage deletion mutants.

Table 4. Effect of site-directed mutagenesis on restoration of cytopathogenicity of pWE2200 and pWE2100

	Strain	Cleavage site	E2	E1	Viability
	pWE2000	RRPKR			Yes
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	pWE2200	R			No
	pWE2221	R		G(2)	No
	pWE2222	R	K(181)	G(2)	Yes
15	pWE2223	R	K(181)		Yes
	pWE2100				No
•	pWE2102		K(182)		Yes

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Although the cleavage deletion mutations in pWE2200 and pWE2100 differed by a single amino acid, the results indicate that the mutations at E2 residues 181 and 182 are both capable of restoring viability of the virus and appear to be equivalent.

EXAMPLE 3 Attenuation of WE2000 virus.

C57/BL6 mice are uniformly susceptible to lethal

challenge by western equine encephalitis virus until
approximately 9 weeks of age. Subcutaneous
inoculation of five- or eight-week-old C57/BL6 female
mice with CBA/87 or WE2000 viruses routinely results
in lethal encephalitis after 8-9 days (Table 2). As

noted previously, the WE2000 is slightly less virulent
than the CBA/87 parent. The virulence of the progeny
virus derived from pWE2100 and pWE2200 infectious
clones were determined by subcutaneous inoculation of
C57BL6 mice. In each instance, the viruses were

significantly attenuated compared to virus produced from parent WE2000 clone. However, infection of mice with increasing doses of WE2215, WE2219 or WE2220, resulted in sporadic deaths with slightly extended periods prior to death compared to the parental virus (Table 5). These results indicated that deletion of only four amino acids from the cleavage site was inadequately attenuating, unlike the results obtained with VEE virus by Davis et al. (1995) and the viruses derived from WE2200 were not characterized further.

Table 5. C57BL6 Mice Inoculated Subcutaneously with WE2200 Cleavage Deletion Mutants

<u>Virus</u>	Mor-	Mean	Prechallenge	Chal	Post Cha	llenge
<u>Strain</u> Dose	taliy	day to Death	ELISA Neut	leng e	ELISA	Neut
CBA/87						
10³	10/10	9.1				
105	10/10	7.7				
WE2000						
103	10/10	9.2				
105	10/10	9.1				·
WE2215						
103	1/10	12	504 (3/9) <20(9/9)	8/9	8300	761
105	1/10	14	100 (6/9) 253 (7/9) <20(9/9)	9/9	12800	403
			100 (2/2)	5,5	12000	403
WE2219						
10³	0/10	-	283(2/8) <20(10/10)	6/10	14368	2281
105	2 /1 0	* •	100 (6/8)			
10	2/10	12	200 (4/8) <20(8/8) 100 (4/8)	8/8	11738	1522
WE2220						
103	1/10	11	606 (5/9) 20(3/9) 100 (4/9) <20 (6/9)	5/9	14703	1280
105	2/10	14	800 (7/8) 20 (2/8)	8/8	9051	761
			100 (1/8) <20 (6/8)			

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When C57BL6 mice were inoculated subcutaneously with the uncloned WE2100 progeny virus, there were no deaths at any dilution, even at doses of 10° PFU per mouse (Table 6). However, some of those mice inoculated subcutaneously with 10⁵ PFU or less remained susceptible to a lethal challenge with the virulent CBA/87 virus.

10 Table 6. C57 Black/6 Mice Inoculated Subcutaneously with WEE Virus Strain WE2100

	strain	Morta:	_		_	Challe	_
Do	se	()	to De	ath	(୫)	
WE2100							
10^{3}		0/10	(0)		-	4/10	(40)
104		0/10	(0)		-	7/10	(70)
10 ⁵		0/10	(0)		_	8/10	(80)
10°		0/10	(0)		-	10/10	(100)
10 ⁷		0/10	(0)	-	-	10/10	(100)

Expressed as animals dying/animals tested
 Expressed as animals surving/animals tested

WE2102 virus was demonstrated to be highly attenuated and killed only two of twenty mice when inoculated subcutaneously with the highest dosage of virus (10° PFU). All mice were challenged 3 weeks later with 10° PFU of CBA/87 virus. Mice previously immunized with 10° PFU or more of WE2102 survived with no noticeable symptoms. Thus, an effective immunizing dose of WE2102 is at least 100 fold less than that required to kill C57BL6 mice. These results further indicate that the Glu to Lys change at position 182 of the E2 glycoprotein is responsible for restoring viability to viruses containing a deletion of the furin cleavage site in the WEE glycoproteins and that WE2102 virus is an effective attenuated vaccine virus.

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Table 7. Subcutaneous Inoculation of C57BL/6 Mice with recombinant WEE virus strain 2102 confers protection against a lethal challenge

Virus Strain Dose WE2100	Mortality ¹ (%)	Mean Day to Death	Challenge ² (%)
10 ³ 10 ⁵ 10 ⁷	0/10 (0) 1/10 (10) 0/10 (0)	16	3/10 (30) 9/9 (100) 10/10 (100)
WE2102 103 105 107	0/20 (0) 0/20 (0) 2/20 (0)	- - 14	9/20 (45) 18/20 (90) 18/18 (100)
WE2000	10/10 (100)	9.2	

- 1. Expressed as animals dying/animals tested
- 2. Expressed as animals surving/animals tested

10 Attenuated VEE IE EXAMPLE 4

Sequence Analysis. Genomic RNA from VEE IE Strain 68U201 was isolated and reverse transcription of the genomic RNA followed by polymerase chain reaction (RT-PCR) was used to generate cDNA of the virus. Initial sequencing of the strain 68U201 genome employed oligonucleotide primers based on existing VEE IE sequence and VEE IA/B sequence. After a partial sequence of strain 68U201 was determined, oligonucleotides specific to the strain 68U201 sequence were used to obtain a complete sequence of the strain 68U201 viral genome. The exact 5' end of the genome was determined by PCR/RACE technique (Frohman et al. [1988] Proc. Natl. Acad. Sci. U.S.A. 85:8998-9002). The entire strain 68U201 viral genomes consists of 11,464

nucleotides, excluding the poly-A sequence (Oberste et al. [1996] Virology 219: 314-320).

EXAMPLE 5

5 Construction of Full-length, Live Clones

Using oligonucleotides specific to the VEE IE strain 68U201 sequence, RT-PCR of strain 68U201 viral RNA was carried out to generate numerous cDNA fragments that were subsequently cloned. These clones 10 were used to assemble a full-length cDNA of strain 68U201 in a plasmid situated such that the cDNA could be precisely transcribed in an in vitro transcription reaction employing T7 polymerase. The first nucleotide downstream of the T7 promoter is a G 15 followed by a cDNA encoding the entire strain 68U201 genome, including poly-A sequence. For the purposes of run off in vitro transcription, a unique endonuclease restriction site (NotI) follows the poly-A sequence (Figure 4). DNA sequences encoding the T7 20 promoter and the strain 68U201 genome were cloned into a suitable plasmid for propagation and selection in E. coli. Oligonucleotides relevant to construction of full-length infectious clones are shown in Table 8.

25 <u>Table 8</u>

0077 CTAAGAGGGCCCCTATATC (SEQ ID NO:16) GCGGAATTCTAATACGACTCACTATAGATGGGCGCGCATGAGAG 0111 (SEQ ID NO:17) 30 0113 TGACCGCGGGACCTCTGTCCAC (SEQ ID NO:18) 0126 AAGTGCATCGATTCAGCG (SEQ ID NO:19) 0136 CTGAAATGTCCAGGATCCACGGAGGAGCTG (SEQ ID NO:20) 0137 CAGCTCCTCCGTGGATCCTGGACATTTCAG (SEQ ID NO:21) 0140 GACTGCGGCCGCTTTTTTTTTTTTGAAATATTAAAAACAAAATCC 35 (SEQ ID NO:22) 0220 CGAGAATCGATGCACTTCAGCC (SEQ ID NO:23)

Descriptions of oligonucleotides are as follows: 0077 introduces an ApaI endonuclease restriction site along with a serine to proline mutation within the coding region of nsP4. 0111 encodes an EcoRI endonuclease restriction site followed by the T7 5 promoter, a single G, and the first 18 nucleotides of the VEE IE genome. 0126 introduces a ClaI endonuclease restriction site within the coding region of nsP3 and was used in combination with 0220. 0113 10 was used to remove an Apal endonuclease restriction site within the structural genes. 0140 encodes a unique NotI endonuclease restriction site followed by a 12 T's, and the reverse complement of the last 21 nucleotides of the VEE IE. A NotI site at the end of 15 the cDNA encoding the VEE IE genome allowed for runoff transcription after digestion of the plasmid with NotI. 0220 introduces a ClaI endonuclease restriction site within the coding region of nsP3 and was used in combination with 0126. Minor alterations of the nucleotide sequence using 0077, 0113, 0126, and 0220 20 facilitated assembly of the full-length clones and allowed for rapid diagnostic analysis of virus generated from these clones by RT/PCR methods.

The full-length clone obtained, pIE1006, was transcribed in vitro using T7 polymerase of NotI-25 linearized plasmid and the RNA transfected into BHK-21 cells (Figure 4). The phenotype of the resulting virus was markedly different from the parent virus (strain 68U201) from which the cDNA was derived. The 30 virus derived from pIE1006 in vitro transcribed RNA, VIE1006, gave rise to small plaques upon infection of target monolayers (Table 9). In attempts to recover the phenotypic characteristics of strain 68U201, regions of the pIE1006 clone were replaced with cDNA generated by RT-PCR from strain 68U201 RNA. Three

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subsequent full-length, live clones were constructed by replacing pIE1006 sequences with 928, 4492, and 8130 nucleotides generated by RT-PCR of strain 68U201 RNA. These clones were designated pIE1007, pIE1008 and pIE1009; and viruses derived from these clones were designated VIE1007, VIE1008 and VIE1009, respectively. These clones are shown in Figure 4. The relevant cloning sites and genetic markers are indicated above each clone. Numbers indicate the corresponding nucleotides in the strain 68U201 genome. The T7 promoter is shown. Drawings are not to scale. Shaded areas represent regions of pIE1006 that were replaced or mutated to generate new clones. The characteristics of these viruses are described below.

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EXAMPLE 6

Assessment of Viruses Derived from Various

Clones. The viruses VIE1006, VIE1007, VIE1008 and

VIE1009 were derived from plasmids pIE1006, pIE1007,

pIE1008, and pIE1009, respectively. Analysis of virus
derived from these clones by plaque assay is shown in

Table 9. Plaque size was determined by infection of

Vero cell monolayers followed by agarose overlay.

Analysis of virus derived from these clones (Table 9) by plaque assay indicated that VIE1007 gave the same plaque morphology as VIE1006 and therefore was not studied further. The VIE1008 produced larger plaques in comparison to VIE1006, and the VIE1009 virus gave rise to the largest sized plaques of the four viruses tested. Plaques generated by VIE1009 were similar to those produced by the parental virus, strain 68U201.

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Table 9: In vitro analysis of virus derived from molecular clones of VEE IE strain 68U201

5	Virus Strain¹	Plaque Size
	VIE1006	small
	VIE1007	small
	VIE1008	medium
10	VIE1009	large
	VEE IE 68	3U201 large

EXAMPLE 7

15 In Vivo studies of VEE IE Vaccine Candidates. The distinct plaque morphologies of VIE1006, VIE1008, and VIE1009 suggested that these viruses may behave differently in vivo. To assess the relative virulence of the cloned derivatives of strain 68U201, mice were 20 initially inoculated with subcutaneously VIE1006, VIE1008, and VIE1009 viruses (Table 10). Mice infected with VIE1006 and VIE1008 were not adversely affected by these viruses as assayed by the number of mice surviving the infection. VIE1009 proved to be as 25 virulent as strain 68U201, causing death in all of the animals infected (Table 10). Immunogenicity of the different viruses inferred by the demonstration of a protective immune response and was determined by back challenge of surviving animals in the virulence assay 30 with approximately 10^4 pfu of the virulent, parental virus, strain 68U201. Back challenge was performed four weeks after the initial inoculation.

Table 10: In vivo analysis of virus derived from molecular clones of VEE IE strain 68U201

5	Virus Strain¹	Mortality ²	Challenge ³
	VIE1006	0/10	10/10
	VIE1008	0/10	10/10
	VIE1009	10/10	nd
10	VEE IE 68U201	nd	1/10

- ¹ Initial inoculation with approximately 10⁴ PFU of each virus with the exception of VEE IE 68U201 which were left untreated until challenge phase of experiment.
- ² Expressed as animals dying/animals tested.
- ³ Expressed as animals surviving/animals tested. All animals were challenged with approximately 10⁴ PFU of VEE IE strain 68U201 four weeks after the initial

20 inoculation.

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EXAMPLE 8

Construction of a Full-length, Molecularly Attenuated VEE IE Clone. With the availability of a 25 full-length virulent clone, specific attenuating mutations were introduced into the structural genes of the virus by site-directed mutagenesis. A deletion mutation was used instead of a point mutation because of the inability of viral RNA replication to repair 30 such mutations. The glycoproteins of VEE IE are produced as a poly-protein precursor, PE2. The junction between the E3 and E2 proteins is cleaved by a furin-like cellular protease. The amino acid sequence of the presumed furin-like protease cleavage site of strain 68U201 is RGKR. The nucleotides encoding these four amino acids of the furin-like cleavage site between the E3 and E2

proteins were deleted from the pIE1009 clone and the resulting "cleavage deletion" clone was designated pIE1100. The oligonucleotides used to generate the cleavage deletion mutation are shown in Table 8. Oligonucleotide 0136 encodes a cleavage deletion mutation that eliminates the four amino acid furin-like cleavage site found in PE2 and was used in combination with oligonucleotide 0137. 0137 encodes the reverse complement of 0136.

10 Transfection of RNA transcribed from pIE1100 into tissue culture cells required extended incubation periods before viral cytopathic effect became apparent in the cultured cells. This extended incubation period is indicative of transcripts from full-length clones possessing mutations that partially inhibit viral replication. However, such mutations can be suppressed by second-site mutations which arise randomnly via the error prone process of alphavirus replication, resulting in variants with enhanced 20 replication ability. In fact, the culture media from cells transfected with RNA transcribed from pIE1100 contained low titers of infectious virus, VIE1100, which could be amplified to high titer $(4x10^7 \text{ PFU/ml})$ upon subsequent passage. Biochemical analysis showed 25 that this virus had an uncleaved PE2 protein indicating that the cleavage deletion mutation totally prevented proteolytic processing of the surface E2 glycoprotein precursor. The efficacy of VIE1100 virus to serve as a vaccine was evaluated in mice as described below. 30

One specific suppressor mutation is thought to reside at nucleotide 10,181 of the VEE IE genome, a C to U nucleotide substitution resulting in an amino acid change from Serine to Leucine in the El protein positon #57. Ability of this mutation to suppress the

lethal nature of the furin cleavage site deletion mutation was assessed. The cDNA of pIE1100 was mutated by changing nucleotide 10,181 from a C to T. This introduced a serine to leucine change at amino acid number 57 of E1, a mutation found within the stock of VIE1100. The resulting clone was designated pVIE1150. Transfection of RNA from pVIE1150 into BHK-21 cells lead to the production of approximately 106 PFU/ml of virus in the supernatant of the transfected cells at 48 hours post-transfection.

EXAMPLE 9

Vaccine Studies with a Full-length, Molecularly
Attenuated Virus. Balb/C mice were inoculated with

VIE1100 virus at various doses (104, 106, and 107 PFU
per mouse), and was found to be completely attenuated
at a dose 100,000 times higher than that required to
cause lethal disease by the parent virus, strain
68U201 (Table 10). Subsequent challenge of these
animals with virulent strain 68U201 demonstrated that
immunization with VIE1100 virus provided complete
protection from lethal virus challenge (Table 11).

Table 11: Vaccination study with VIE1100

25			
	Virus Strain Dose	Mortality ¹	Challenge ²
	VIE1100		
30	10⁴	10/10	10/10
	10 6	10/10	10/10
	107	10/10	10/10
	Mock Vaccinated	0/10	0/10

- 35 1. Expressed as animals dying/animals tested.
 - 2. Expressed as animals surviving/animals tested after inoculation with 10^4 PFU of VEE IE strain 68U201

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EXAMPLE 10

Properties of a WEE-EEE chimeric virus. Based upon similarities of amino acid sequences of the carboxy terminal portion of NSP-4 and the amino terminal portion of the capsid proteins of WEE, EEE and Sindbis virus, it has been suggested that WEE virus arose by recombination between EEE and a Sindbis virus ancestor (Hahn et al. [1988] Proc. Natl. Acad. Sci. U.S.A. 85: 5997-6001). The capsid genes of both WEE and EEE contain a highly conserved sequence with a 10 unique Bln I site 76 nucleotides downstream of the initiation codon. We utilized this site to insert the structural protein sequences of EEE into pWE5-18 in order to construct a full length clone encoding a chimeric virus. Plasmid pMWE-7 is a full length clone 15 consisting of the 5' non-coding sequence, nonstructural genes, 26S promoter and the first 25 codons for the capsid protein of WEE CBA/87 fused to the structural protein genes and 3' non-coding 20 sequence of EEE strain Fla91-4679. Transfection of BHK cells with RNA transcribed from pMWE-7 resulted in complete destruction of the monolayer and release of high yields of virus. SDS-polyacrylamide gel electrophoresis of the purified virus demonstrated 25 that the virus is composed of polypeptides which comigrate with those of EEE and not of WEE, indicating that the virus is a chimera (Figure 3).

Injection of mice with the MWE chimeric virus killed mice only sporadically, suggesting that fusion of the sequences of the two viruses resulted in significant attenuation compared to the parent WE2000 virus (Table 12). Mice immunized with MWE-7 developed significant neutralizing antibody titers and resisted a lethal EEE Fla91-1467 challenge when immunized with greater than 10⁵ plaque forming units of

MWE-7 virus. The neutralizing antibody response after challenge was not significantly elevated indicating that the immunization with the chimeric virus effectively prevented infection by a lethal EEE challenge. Therefore, a chimeric virus derived by combining the structural protein genes of EEE with the non-structural proteins genes of WEE may serve as a safe, effective approach to development of a vaccine for EEE virus.

Table 12. Immunization of C57BL6 Mice with an WEE/EEE Chimeric Virus Confers Protection Against a Lethal EEE Challenge

Virus Strain Dose WE2000	Mortality ¹	Mean Day to Death	Challenge ²
10 ⁵	10/10 (100)	8.5	
MWE7			
10³	0/10 (0)		6/10 (60)
10 ⁵	1/10 (10)	8.0	9/9 (100)
10 ⁷	$0/10 (0)^3$	0.0	, , , , , , , , , , , , , , , , , , , ,
10	0/10 (0/		9/9 (100)
Saline Control	0/10 (0)		0/10 (0)

- 15 1. Expressed as animals dying/animals tested
 - 2. Expressed as animals surving/animals tested
 - 3. One animal died during pre-challenge bleed

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Example 12 VEE IIIA Chimera

Construction of IAB-IIIA cDNA Chimeric Clone (pV3A-1000)

BeAn8 (wild type Mucambo or VEE IIIA) cDNA was amplified with primers 128 and 140 (Table 13) to generate a PCR fragment of the entire 26S region of BeAn8. Primer 128 incorporated an ApaI site just upstream of the BeAn8 26S promoter to facilitate cloning into pV3000. Primer 140 introduced a NotI

site just downstream of the poly(A) tract which would also facilitate cloning into pV3000 as well as enable run-off transcription. This 128/140 PCR fragment was cloned into pBluescript SK+ vector using the ApaI and NotI sites and was termed pMUC-1000.

To replace the BeAn8 3' nontranslated region (3'NTR) in pMUC-1000 with the corresponding region in pV3000, an EcoRI site was introduced immediately downstream of the E1 stop codon in both pMUC-1000 and pV3000 using the primer pairs 013/227 and 225/226, 10 respectively. The 013 primer is a universal primer found in the pBluescript SK+ vector and the 227 primer was designed to introduce an EcoRI site in pMUC-1000. The 225 primer introduced an EcoRI site in pV3000 and the 226 primer is located in the vector, TotoX. The 013/227 and 225/226 PCR fragments were separately cloned into the pBluescript SK+ vector producing pMUC-1200 and pV3nt-1000, respectively. The 3' NTR encoded in pV3nt-1000 was shuttled into pMUC-1200 using the 20 EcoRI site and the NotI site to produce pMUC-1300.

The full-length chimeric cDNA clone, pV3A-1000, was constructed by shuttling the structural genes encoded in pMUC-1300 into the pV3000 nonstructural domain using the ApaI site and NotI site.

Table 13. Primers for construction of the pV3A-1000

013 -- AACAGCTATGACCATG (SEQ ID NO:24)

128 -- CTGAGAGGGGCCCCAGTAAC (SEQ ID NO:25)

30 140 -- GACTGCGGCCGCTTTTTTTTTTTTGAAATATTAAAAA (SEQ ID NO: 26)

225 -- CCAGAAACATAATTGAATTCAGCAGCAATTG (SEQ ID NO:27)

226 -- CTTTATCCGCCTCCATCC (SEQ ID NO:28)

227 -- CCAATCGCTGCTGAATTCTAATTATGTTTCTG (SEQ ID NO:29)

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Five week old C57BL6 mice were used in a challenge study (Table 14). All mice immunized with

10⁷ PFU of pV3A-1000 produced neutralizing antibodies. Mice were challenged with 10⁷ PFU of 900807 (a Mucambo virus from Trinidad). The geometric mean neutralizing antibody titer in mice inoculated with 10⁷, 10⁵, and 10³ PFU of pV3A-1000 was 9800, 8800, and 7000 respectively.

	<u> Table 14.</u>	Results f	rom chall	enge studies	1
10		Fraction	Fraction of mice	Average days to onset of	Average days to death
15	Immunized with pV3A-1000	0/20	0/20	N/A	N/A
20	Unvac- : cinated	19/20	12/20	6.2	10.5

SEQUENCE LISTING

	(I) GENERAL INFORMATION:
5	(i) APPLICANT: Michael D. Parker Jonathan F. Smith Bruce Crise
	Mark Steve Oberste
10	Shannon Schmura
	(ii) TITLE OF INVENTION: Live Attenuated virus
	vaccines for western equine encephalitis virus,
	eastern equine encephalitis virus, and venezuelan
1 F	equine encephalitis virus IE and IIIA variants
15	
	(iii) NUMBER OF SEQUENCES:29
	(iv) CORRESPONDENCE ADDRESS:
20	(A) ADDRESSEE: John Moran
20	(B) STREET: USA MRMC - MRMC-JA (C) CITY: FORT DETRICK, FREDERICK
	(D) STATE: MARYLAND
	(E) COUNTRY: USA (F) ZIP: 21702-5012
25	(r) BIF: Z1/02-3012
	(v) COMPUTER READABLE FORM:
	(A) MEDIUM TYPE: Floppy disk
	(B) COMPUTER: Apple Macintosh (C) OPERATING SYSTEM: Macintosh 7.5
30	(D) SOFTWARE: Microsoft Word 6.0
	<pre>(vi) CURRENT APPLICATION DATA: (A) APPLICATION NUMBER:</pre>
	(B) FILING DATE:
35	(C) CLASSIFICATION:
	<pre>(vii) PRIOR APPLICATION DATA: (A) APPLICATION NUMBER: Provisional</pre>
	Application 60/047162,
40	(B) FILING DATE: May 20, 1997
	(4)
	(A) APPLICATION NUMBER: Provisional
	Application 60/053,652

	(B) FILING DATE: July 24, 1997
5	<pre>(viii) ATTORNEY/AGENT INFORMATION: (A) NAME: Moran, John (B) REGISTRATION NUMBER: 26,313 (C) REFERENCE/DOCKET NUMBER:</pre>
10	<pre>(ix) TELECOMMUNICATION INFORMATION (A) TELEPHONE: (301) 619-2065 (B) TELEFAX: (301) 619-7714</pre>
	(2) INFORMATION FOR SEQ ID NO:1:
15	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 11492 base pairs (B) TYPE: Nucleic acid (C) STRANDEDNESS:Double (D) TOPOLOGY: Linear
20	(ii) SEQUENCE DESCRIPTION: SEQ ID NO:1: 1 ATAGGGCATG GTATAGAGGC ACCTACCCTA CAAACAAATC
	GATCCAATAT
25	51 GGAAAGAATT CACGTTGACT TAGACGCTGA CAGCCCATAT
	101 TACAGCGGAC GTTTCCACAA TTTGAGATCG AAGCAAGGCA
30	151 AATGACCATG CCAATGCCAG AGCGTTTTCG CATGTGGCAA
35	201 TGAGAGCGAA GTCGACCGGG ACCAAGTTAT CTTGGACATT GGAAGTGCGC
	251 CCGTCAGACA TGCACATTCC AACCACCGCT ATCATTGTAT
40	301 ATAAGCGCTG AAGACCCGGA CAGACTACAG CGGTATGCAG AAAGACTTAA
	351 GAGAAGTGAC ATGTACCGAC AAGAATATAG CCTCTNAAGG

401 GCTGGAAGTC ATGTCCACAC CAGACGCAGA GACTCCATCT

451 ACACAGACGC CACGTGTAGG TACTTTGGAA GTGTANGCAG

50

45

CTGTGTATGC

TATACCAAGA

TGACTCATTT

	GCGCTTAAAG	GICCATGCAC	CGACATCAAT	CTACCACCAC
5	551 GAGTTAGGAC TTTTATGTAC	AATTTACTGG	ATAGGTTTTG	ACACGACCC
	601 AAAAACATGG GGGCCGACGA	CAGGTTCCTA	CCCTACTTAC	AACACAAACT
10	651 GAGAGTATTG GATCTTCAGG	GAAGCACGTA	ACATTGGCCT	CGGTAACTCA
15	701 AGAGCAGACT GCTCCAACCT	TGGAAAACTT	TCAATCCTTA	GGAAGAAGAG
13	751 ACTGATAAGA CAGAGGATAG	TCATATTCTC	GGTTGGTTCA	ACAATCTACA
20	801 ATCACTGTTA CTGAAAGGAA	CGTAGCTGGC	ATCTTCCAAA	CGTGTTCCAC
	851 AGTCTNACTT TGAAGGGTAT	CACAGGTAGA	TGTGGGACCA	TTGTCAGCTG
25	901 GTCATCAAAA AAGTTGAGAA	AGATAACGAT	CAGCCCAGGA	CTATACGGTA
30	951 CTTGGCGTCC AAAGTCACAG	ACGATGCATC	GCGAGGGTTT	CTTGAGTTGC
50	1001 ATACGCTGCG GTATGTACCA	CGGCGAGAGG	GTTTCTTTTG	CTGTGTGTAC
35	1051 GCCACACTTT CGGATTGTCG	GCGATCAGAT	GACAGGGAGG	GCTCAACCAA
	1101 TCAATGGTAG CTATCTATTA	GACGCAAAGA	AATACTNACA	CAATGCAGAA
10	1151 CCAGTGGTCG ATCGTGCCGA	CCCAGGCGTT	TTCCAGGTGG	GCGCGTGNAC
15	1201 CTTGGACGAC CTTACTATGG	GAGAAGGAGC	TAGGGGTGCG	GGAGCGCACT
ŧJ	1251 GCTGCTGCTG CTACAAGAAG	GGCTTTCAAG	ACCCAGAAAA	TTACATCCAT
	1301 CCTGGTACGC	АААСАААТТА	AGAAAGTACC	ጥርርርርጥርጥጥጥ

	1351 GTGATTCCGA CGCCTTACCA GCCACGCGGG GGCTCGA TGGGCCTTCC	ATA
5	1401 GCCGTNAGGC TCAAGCTGCT GCTTGAACCA ACTGTCA	AAC
	1451 TATTACAATG GCCGATGTGG AGCACCTGCG TGGCTTA CAAGAAGCTG	CAG
10	1501 AAGAAGTGGC TGCAGCGGGA AGAGATCAGA GAAGCCCCCACCCTTGCT	TGC
15	1551 CCCTGAAATA GAAAAAGAGA CCGTAGAGGC AGAAGTA	GAC
13	1601 AAGAGGCAGG AGCAGGTAGC GTGGAGACAC CNACGAGATATCAAGGT	GAC
20	1651 AACAAGTTAC CCAGGTGNAA GAGAAGATTG GGTCTTA'	rcc
	1701 ACCCCAGGCG GTTTTANAAT NGTNAAAAAC TGGCGTG	ГАТ
25	1751 GCGGAACAAG TACTGGTAAT GACTCACAAA GGCAGGGGGGGGGATACAA	CCG
30	1801 AGTCGAGCCA TACCACGGTA NGGTCATTGT ACCCAGAIGACGGCGGGT	AGG
30	1851 CCCTGTTCAA GACTTTCAGG CACTGAGTGA GAGCGCCAATCGTTTTCA	4CG
35	1901 ACGAGAGGGA GTTCGTAAAC CAGATATTTT GCACCCACCGCAAGCKTT	CAT
	1951 TCAACTATAG TGAGTCGCTA TTACACTGAC GAAGAGTA	¥СТ
40	2001 AAAGACTCAG GACGCAGACT CAGAATACGT CTTTGACAGACGCACGAA	\TT
4 =	2051 AGTGTGTTAA GCGAGAAGAC GCAGGTCCCT TGTGCCTA	ł A C
45	2101 GTAGATCCAC CATTTCACGA GTTTGCGTAC GAGAGTCTAGACACGACC	rca
50	2151 AGCAGCACCT CACAAAGTCC CAACCATCGG AGTCTATC	3GA

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	2201 CAGGTAAATC TGGGATCATC AAAAGCGCTG TGACTAAGAA AGATCTGGTT
5	2251 GTGAGTGCGA AGAAGGAAAA CTGCGCAGAA ATTATCAGGG ATGTAAGGAG
	2301 GATGAGACGT ATGGATGTTG CTGCTAGGAC TGTTGATTCA GTGCTTCTAA
10	2351 ATGGGGTTAA GCACCCCGTT AACACCCTGT ACATTGATGA GGCATTTGCC
15	2401 TGCCATGCAG GGACGCTGCT GGCACTGATT GCCATCGTCA AACCTAAGAA
15	2451 AGTGGTATTG TGCGGGGACC CAAAACAATG CGGCTTCTTT AACATGATGT
20	2501 GCCTGAAAGT ACATTTTAAC CATGACATAT GCACTGAGGT GTACCACAAA
	2551 AGCATCTCTA GGAGGTGCAC ACAGACTGTA ACCGCCATTG
25	2601 CTTCTACGAC AAGCGAATGA AGACGGTTAA CCCATGTGCT GACAAAATCA
	2651 TCATAGATAC CACAGGGACC ACAAAGCCGC ACAAAGATGA
30	2701 ACCTGTTTCA GAGGATGGGT GAAACAGCTA CAGATTGACT ATAAAAATCA
35	2751 TGAAATCATG ACTGCGGCTG CATCGCAAGG ACTTACGCGG AAAGGCGTTT
	2801 ATGCTGTCAG GTACAAAGTC AACGAGAATC CACTCTACTC GCAGACTTCT
40	2851 GAGCACGTGA ACGTGTTAGT TACACGCACA GAAAAACGCA TTGTCTGGAA
	2901 GACGCTAGCT GGTGACCCCT GGATAAAGAC ACTTACAGCT AAATACTCCG
45	2951 GGGATTTCAC GGCTTCATTG GACGACTGGC AACGCGAACA
50	3001 ATGGCACGCG TTCTTGATAA GCCGCAGACA GCTGATGTGT TCCAGAATAA

	GCCACGGCCA	TGCTGGGCGA	AGGCTTTAGA	GCCAGTCTTC
5	3101 ACATTGTGCT ATTCAAGCAT	GACGAGACAG	CAGTGGGAGA	CGTTGCACCC
	3151 GACAGAGCGT GCACCAGGTT	ACTCACCTGA	AATGGCACTG	AACTTCTTTT
10	3201 CTTTGGTGTA ACCGTCGCAC	GACCTGGACA	GTGGGTTGTT	TTCCGCTCCT
15	3251 TTACTTACAG GAACATGTAT	GGATCAGCAC	TGGGACAACT	CGCCAGGGAA
13	3301 GGGCTTAATA ATCCGTGCAT	GAGAGGTAGC	AAAGGAGCTG	TCACGGCGAT
20	3351 CACAAAAGCG AATAATACCA	GTTGACACAG	GCAGGGTAGC	TGATATAAGG
	3401 TCAAGGACTA TCGCCGGTTG	CTCTCCAACA	ATTAATGTGG	TTCCATTAAA
25	3451 CCCCACTCGC CTGATCACAG	TGATCGTTGA	CCACAAAGGA	CAGGGTACAA
30	3501 TGGATTCCTA ATCGGCGATC	TCTAAGATGA	ATGGCAAATC	TGTGTTGGTG
	3551 CTATCAGCAT ATTGCCCACT	TCCTGGGAAG	AAAGTAGAGT	CCATGGGTCC
35	3601 AATACCATCA (ATGTCGGTAA	GGTGTGATCT	CGATTTGGGA	ATACCTAGCC
	3651 ATATGACATT AACCATCACT	ATATTTGTCA	ATGTTAGGAC	CCCATACAAG
40	3701 ACCAACAGTG (AACGTGTAAG	CGAGGATCAC	GCTATCCACC	ACAGCATGCT
45	3751 GCTGTCCACC ATAGGGTATGG	ACCTGAACAC	TGGCGGAACA	TGTGTGGCCA
	3801 ACTTGCTGAT GCTCGCTCAT	CGCGCAACCG	AGAATATCAT	CACTGCGGTG
50	3851 TTAGGTTTAC AAATACTGAG	CCGTGTCTGT	CAGCCTAAGA	ACACTGCCGA

3901 GTTCTCTTCG TGTTCTTCGG CAAGGACAAC GGCAACCACA CACATGACCA 3951 GGACAGACTC GGTGTAGTGC TTGACAACAT CTACCAAGGG TCAACCAGGT 4001 ACGAGGCAGG GAGAGCTCCA GCGTACAGAG TGATCAGAGG TGACATTAGC 10 4051 AAGAGCGCTG ACCAAGCTAT CGTTAATGCT GCTAATAGCA AAGGTCAACC 4101 AGGTTCCGGA GTGTGCGGTG CACTGTACCG AAAATGGCCG GCTGCTTTTG 15 4151 ATAGACAGCC AATAGCTGTC GGGACGGCTA GACTTGTGAA GCACGAACCG 4201 CTCATCATAC ATGCTGTAGG ACCGAATTTT TCTAAGATGC 20 CGGAACGAGA 4251 GGGCGACCTT AAGCTCGCAG CTGCCTACAT GAGCATAGCG TCCATCGTCA 25 4301 ACGCTGAGCG GATTACTAAA ATATCAGTAC CGCTACTGTC **AACTGGCATC** 4351 TATTCTGGTG GCAAAGATCG AGTGATGCAA TCATTGCATC ACTTGTTCAC 30 4401 TGCTTTCGAC ACTACGGATG CCGATGTCAC CATATATTGC TTGGATAAAC 4451 AATGGGAGAC CAGGATAACC GAGGCCATTC ACCGCAAAGA 35 AAGCGTCGAA 4501 ATACTGGATG ATGATGACAA GCCAGTAGAC ATCGACTTGG TCAGGGTCCA 40 4551 CCCAAACAGC TCTTTGGCAG GCAGACCAGG TTACTCCGTC AATGAGGGCA 4601 AGCTGTATTC ATACCTGGAA GGTACACGAT TCCATCAGAC CGCCAAGGAC 45 4651 ATTGCCGAAA TCCATGCAAT GTGGCCCAAC AAATCTGAGG CTAATGAGCA 4701 GATTTGCTTG TACATCCTGG GGGAGAGTAT GTCCAGCATC 50 CGCTCCAAAT

4751 GCCCAGTAGA GGAGTCAGAG GCGTCTGCTC CACCTCACAC ACTTCCATGC 4801 CTGTGTAATT ACGCTATGAC GGCTGAGCGC GTATACAGGT TGCGCTCTGC 4851 GAAGAAAGAA CAGTTCGCCG TATGCTCATC ATTCCCGTTG CCGAAGTACA 10 4901 GGATCACAGG CGTGCAGAAG CTACAGTGCA GCAAACCAGT CCTGTTCTCA 4951 GGCGTCGTAC CACCGGCTGT ACACCCCAGG AAGTACGCGG AAATAATTCT 15 5001 AGAAACGCCA CCATCGCCAA CAACGACAAC CGTAATATGT GAACCAACTG 5051 TGCCAGAACG TATACCCAGT CCGGTGATTT CTAGAGCAGC 20 AAGTGCGGAA 5101 TCACTGCTAT CGTTTGGCGG CGTCTCGTTC TCTAGCTCTG CCACACGCTC 25 5151 GTCAACTGCC TGGAGCGACT ATGACAGGCG GTTTGTGGTT ACAGCTGACG 5201 TGCATCAAGC GGACACGTCT ACGTGGAGCA TCCCTAGCGC TCCTGACTTG 30 5251 GACGTCCAGC TGCCTTCTGA CGATACTGAT TCCCACTGGA GTATTCCGAG 5301 TGCATCAGGC TTTGAAGTGA GAACACCGTC TGTACAGGAC 35 CTAACTGCAG 5351 AGTGTGCGAG GCCTCGTGGA CTGGCCGAAA TAATGCAAGA CTTCAATACG 40 5401 GCTCCTTTCC AGTTTCTTTC GGACCACAGA CCAGTACCAG CACCACGGAG 5451 ACGCCCCATC CCATCACCTA GATCGACGGT TTCCGCACCT CCAGTTCCAA 45 5501 AGCCACGCAG AACTAAGTAC CAACAACCAC CAGGAGTCGC TAGGGCGATC 5551 TCAGAAGCGG AGCTGGACGA GTACATCCGT CAACATTCCA 50 ATTGACGGTA

5601 TGAAGCGGAG CGTATATTTC TCATCGAAAC AGGCCAGGTC ACCTTCAACA 5651 GAAATCAGTA CGTCAATGTA AACTACAAGA CCCTATATTG GAACGGGCCG 5701 TCCATGAGAA GTATTAGCCC CGCGCCTCGA TCTCGAAAGA GAGAAGATGT 5751 TACAGAAGAA ACTGCAATTA TGCGCCTCNT GAAGGAAATA 10 GAAGCAGGTA 5801 TCAATCACGA AAAGTANGAA AATATGAAAG CAATTACAGC GGAGCGACTC 15 5851 ATTTCTGGAT TGGGCACATA TCTATCATCA GAAGTGAATC CTGTCGAGTG 5901 TTACAGAGTC AATTATCCTG TACCAATCTA CTCGTCAACG 20 GTAGTTAATA 5951 GGTTTTCATC AGCAGAAGTG GCCGTCAAAG TTTGCAACTT AGTCATCCAA 25 6001 GAGAATTACC CTACAGTAGC CAGTTATTGC ATAACAGATG AATACGATGC 6051 GTATCTTGAC ATGGTGTACG GCGCTCGTGC TGTTAGATAC AGCGCCTTTG 30 6101 TCCGCTNACT GAGAAGCTAC CCAAAGAAGC ATAGCTACTT GCAGCCAGAG 6151 ATAAGATCAG CTGTCCCATC ACCTATACAG AATACATTAC 35 AAAATGTATT 6201 GCTGCAGCCA CTAAAAGGAA CTGCAACGTT ACCCAAATGC GAGAACTACC 6251 TGTTTTATTC GGCGGCATTC AACGTTGAAT GTTTCAAGAA 40 ATACGCATGC 6301 AATGATGAGT ATTGGGATAC CTTTCGCGAT AACCCTATTC GGCTAACTAC 45 6351 AGAGAACGTT ACGCAATACG TGACAAAGCT GAAAGGGCCG AAAGCAGCAG 6401 CATTATTCGC AAAAACTCAC AACCTAAAAC CGTTGCAGGA 50 GATACCAATG

GGTGAAGGCC

6451 GACGAATTTG TCATGGTCTN AAAAGAGATG TCAAAGTTAC TCCCGGCACA 6501 AAACATACAG AGGAGCGGCC TAAGGTGCAG GTTATTCAGG CTGCAGATCC 6551 TCTCTGTACC GCTTACCTTT GCGGGATCCA TCGAGAATGT CCGTAGACTG 10 6601 AATGCTGTGC TTCTGCCGAA TATCCATACT CTCTTCGACA TGTCAGCGGA 6651 AGATTTTGAT GCGATTATTG CTGAACATTT CCACCACGGC GACCCAGTAT 15 6701 TGGAAACGGA CATCGCGTCG TTTGATAAAA GCGAAGACGA CGCTATCGCC 6751 ATTTCGGCGT TGATGATCCT CGAGGACTTA GGCGTCGACC 20 AACCGCTCTT 6801 AGATTTGATA GAGGCGGCGT TCGGCAATAT CACATCTGTG CACCTACCTA 6851 CAGGAACGAG GTTCAAATTT GGTGCCATGA TGAAATCCGG 25 TATGTTCTTA 6901 ACGCTGTTTG TCAACACACT AGTCAATATC ATGATTGCTA **GCAGAGTACT** 30 6951 ACGTGAACGG TTAACCACGT CAGAGTGCGC GGCCTTATCG GCGACAATAA 7001 CATAGTGCAC GGTGTCGCTC CGACAACTTG ATGGCGGAGA 35 GATGCGCCAC 7051 TTGGCTGAAC ATGGAAGTAA AAATTATTGA TGCAGTCATT GGTATCAAAG 40 7101 CACCCTACTT CTGCGGGGAT TCATCCTGGT GGATCAGATA ACGACACAGC 7151 CTGTAGGTCG CAGACCCTCT AAAAAGGCTT TTTAAGCTTG GAAAACCATT 45 7201 GCCAGTCGAT GACACCCAAG ACTGTGACCG TCGCCGGGCA CTGCATGATG 7251 AAGCAATGCG ATGGAACAGA ATTGGAATTA CGGACGAGTT

7301 GTAGAATCCA GATACGAGAT CATACTGGCA GGCCTGATCA TCACGTCCCT 7351 GTCCACGTTA GCCGAAAGCG TTAAGAACTT CAAGAGCATA AGAGGGAACC 7401 CAATCACCCT CTACGGCTGA CCTAAATAGG TGACGTAGTA GACACGCACC 7451 TACCCACCGC CAAAAATGTT TCCATACCCT CAGCTGAACT 10 TTCCACCAGT 7501 TTACCCTACA AATCCGATGG CTTACCGAGA TCCAAACCCT CCTAGGCGCC 15 7551 GCTGGAGGCC GTTTCGGCCC CCGCTGGCTG CTCAAATCGA AGATCTTAGG 7601 AGGTCGATAG CCAACTTAAC TTTCAAACAA CGATCACCTA 20 ATCCGCCGCC 7651 AGGTCCGCCG CCGAAGAAGA AGAAGAGTGC TCCCAAGCCA AAACCTACTC 25 7701 AGCCTAAAAA GAAGAAGCAG CAAGCCAACA AGACGAAACG CAAGCCTAAA 7751 CCAGGGAAAC GACAGCGTAT GTGTATGAAG TTGGAGTCGG ACAAGACATT 30 7801 TCCGATCATG TTGAACGGCC AAGTGAATGG ATACGCTTGC GTTGTCGGAG 7851 GAAGGCTGAT GAAACCACTC CACGTTGAAG GAAAAATCGA 35 TAATGAGCAA 7901 TTAGCGGCCG TGAAATTGAA GAAGGCTAGC ATGTACGACC TGGAGTATGG 7951 CGACGTTCCC CAGAATATGA AATCAGACAC GCTGCAGTAC 40 ACCAGCGACA 8001 AACCACCGGG CTTCTACAAC TGGCACCACG GCGCAGTCCA GTATGAGAAT 45 8051 GGGAGATTCA CCGTACCGCG AGGAGTGGGC GGGAAAGGCG ACAGTGGAAG 8101 ACCGATCCTG GACAACAGAG GCAGAGTTGT GGCTATTGTT 50 CTAGGAGGTG

	8151 CAAACGAGGG GAACCAGAAA	CACGCGTACG	GCGCTTTCAG	TGGTCACTTG
5	8201 GGGGTGACCA GGTCACTAGT	TCAAGGATAC	CCCCGAAGGT	TCTGAACCGT
	8251 TACAGCGCTG GACAAACCAC	TGCGTGCTTT	CGAATGTCAC	ATTCCCTTGC
10	8301 CCGTGTGCTA GCTCGAGGAG	TTCACTGGCG	CCAGAACGAA	CACTCGACGT
15	8351 AACGTCGACA TCTTGAAATG	ATCCAAATTA	CGACACGCTG	CTGGAGAACG
13	8401 TCCATCACGC ACGCTGACCA	CGGCCCAAAC	GAAGCATTAC	CGATGACTTC
20	8451 GTCCCTACCT GCCATGTTTT	GGGGTTCTGC	CCGTATTGCA	GACACTCAGC
	8501 AGCCCAATAA ATGGGTCGAT	AAATTGAGAA	CGTGTGGGAC	GAATCTGATG
25	8551 TAGAATCCAG GGCACTGCAG	GTCTCGGCAC	AATTCGGCTA	CAATCAGGCA
30	8601 ACGTCACCAA TGACATCAAG	GTTCCGGTAC	ATGTCTTACG	ACCACGACCA
50	8651 GAAGACAGTA CATGCCGTCG	TGGAGAAAAT	AGCTATTAGT	ACATCTGGAC
35	8701 TCTTGGCCAC CCAGGTGACA	AAAGGGTACT	TCCTGTTAGC	TCAATGTCCT
	8751 GTGTAACCGT ATGCACCGTG	CAGTATCACG	AGCGGAGCAT	CTGAGAATTC
10	8801 GAGAAAAAGA ACTTGTTCCC	TCAGGAGGAA	GTTTGTCGGT	AGAGAGGAGT
4 =	8851 ACCTGTCCAT CACTTGAAGG	GGAAAGCTGG	TAAAGTGCCA	CGTTTACGAT
15	8901 AGACGTCTGC ACACGCGTAT	CGGATACATA	ACCATGCACA	GGCCAGGCCC
50	8951 AAGTCCTATC AACCACCTTC	TGGAGGAAGC	GTCAGGCGAA	GTGTACATTA

	9001 TGGCAAGAAC GTCACCTACG AATGTAAGTG TGGTGACTAC AGCACAGGTA
5	9051 TTGTGAGCAC GCGAACGAAG ATGAACGGCT GCACTAAAGC AAAACAATGC
	9101 ATTGCCTACA AGCGCGACCA AACGAAATGG GTCTTCAACT CGCCGGATCT
10	9151 TATTAGGCAC ACAGACCACT CAGTGCAAGG TAAACTGCAC ATTCCATTCC
15	9201 GCTTGACACC GACAGTCTGC CCGGTTCCGT TAGCTCACAC GCCTACAGTC
13	9251 ACGAAGTGGT TCAAAGGCAT CACCCTCCAC CTGACTGCAA CGCGACCAAC
20	9301 ATTGCTGACA ACGAGAAAAT TGGGGCTGCG AGCAGACGCA ACAGCAGAAT
	9351 GGATTACAGG GACTACATCC AGGAATTTTT CTGTGGGGCG AGAAGGGCTG
25	9401 GAGTACGTAT GGGGCAACCA TGAACCAGTC AGAGTCTGGG CCCAGGAGTC
30	9451 GGCACCAGGC GACCCACATG GATGGCCGCA TGAGATCATC ATCCACTATT
30	9501 ATCATCGGCA TCCAGTCTAC ACTGTCATTG TGCTGTGGGTGTCGCTCTT
35	9551 GCTATCCTGG TAGGCACTGC ATCGTCAGCA GCTTGTATCG CCAAAGCAAG
	9601 AAGAGACTGC CTGACGCCAT ACGCGCTTGC ACCGAACGCA ACGGTACCCA
40	9651 CAGCATTAGC AGTTTTGTGC TGTATTCGGC CAACCAACGC TGAAACATTT
45	9701 GGAGAAACTT TGAACCATCT GTGGTTTAAC AACCAACCGT TTCTCTGGGC
40	9751 ACAGTTGTGC ATCCCTCTGG CAGCGCTTGT TATTCTGTTC CGCTGCTTTT
50	9801 CATGCTGCAT GCCTTTTTTA TTGGTTGCAG GCGTCTGCCT GGGGAAGGTA

	9851 GACGCCTTCG AACATGCGAC CACTGTGCCA AATGTTCCGG GGATCCCGTA
5	9901 TAAGGCGTTG GTCGAACGTG CAGGTTACGC GCCACTTAAT CTGGAGATTA
	9951 CGGTCGTCTC ATCGGAATTA ACACCCTCAA CTAACAAGGA GTACGTGACC
10	10001 TGCAAATTTC ACACAGTCGT TCCTTCACCA CAAGTTAAAT GCTGCGGGTC
15	10051 CCTCGAGTGT AAGGCATCCT CAAAAGCGGA TTACACATGC CGCGTTTTTG
15	10101 GCGGTGTGTA CCCTTTCATG TGGGGAGGCG CACAGTGCTT CTGTGACAGT
20	10151 GAGAACACAC AACTGAGTGA GGCATACGTC GAGTTCGCTC CAGACTGCAC
	10201 TATAGATCAT GCAGTCGCAC TAAAAGTTCA CACAGCTGCT CTGAAAGTCG
25	10251 GCCTGCGTAT AGTATACGGC AATACCACAG CGCGCCTGGA TACATTCGTC
••	10301 AACGGCGTCA CACCAGGTTC CTCACGGGAC CTGAAGGTCA TAGCAGGGCC
30	10351 GATATCAGCA GCTTTTTCAC CCTTTGACCA TAAGGTCGTC ATTAGAAAGG
35	10401 GGCTTGTTTA CAACTACGAC TTCCCTGAGT ATGGAGCTAT GAACCCAGGA
	10451 GCGTTCGGCG ATATTCAAGC ATCCTCTCTT GATGCCACAG ACATAGTAGC
40	10501 CCGCACCGAC ATACGGCTGC TGAAGCCTTC TGTCAAGAAC ATCCACGTCC
	10551 CCTACACCCA AGCAGTATCA GGGTATGAAA TGTGGAAGAA CAACTCAGGA
45	10601 CGACCCCTGC AAGAAACAGC ACCATTCGGA TGTAAAATTG AAGTGGAGCC
50	10651 TCTGCGAGCG ACTAACTGTG CTTATGGGCA CATCCCTATC

10701 TCCCTGATGC AGCTTTTGTG AGATCATCTG AATCACCAAC AATTTTAGAA 10751 GTCAGCTGCA CAGTAGCAGA CTGCATTTAT TCTGCAGACT TTGGTGGTTC 10801 GCTAACACTA CAGTACAAAG CTAACAGAGA GGGACATTGT CCAGTTCACT 10851 CCCACTCCAC TACAGCTGTT TTGAAGGAAG CGACCACACA 10 TGTGACTGCC 10901 ACAGCCATAA CACTACATTT TAGCACATCG AGCCCACAAG CAAATTTCAT 15 10951 AGTTTCGCTA TGGCGCAAGA AGACCACCTG CAATGCTGAA TGTAAACCAC 11001 CGGCCGACCA CATAATTGGA GAACCACATA AGGTCGACCA 20 AGAATTCCAG 11051 GCGGCAGTTT CCAAAACATC TTGGAACTGG CTGCTTGCAC TGTTTGGGGG 25 11101 AGCATCATCC CTCATTGTTG TAGGACTTAT AGTGTTGGTC TGCAGCTCTA 11151 TGCTTATAAA CACACGTAGA TGACTGAGCG CGGACACTGA CATAGCGGTA 30 11201 AAAACTCGAT GTACTTCCGA GGAAGCGTGG TGCATAATGC CACGCGCCGC 11251 TTGACACTAA AACTCGATGT ATTTCCGAGG AAGCACAGTG 35 CATAATGCTG 11301 TGCAGTGTCA CATTAATCGT ATATTACACT ACATATTAAC AACACTATAT 40 11351 CACTTTTATG AGACTCACTA TGGGTTTCTA ATACACACTA CACACATTTT 11401 ATTTAAAAAC ACTACACACA CTTTATAAAT TCTTTTATAA TTTTTCTTTT 45 11451 GTTTTTTATT TTGTTTTTAA AATTTCAAAA AAAAAAAAA

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٦	. •	TIM OTHER TON	LOK	JEU	$\perp \nu$	INU : Z. :

1	(i)	SECTIENCE	CHARACTERISTICS.

- 5 (A) LENGTH: 11464 base pairs
 - (B) TYPE: Nucleic acid
 - (C) STRANDEDNESS: Double
 - (D) TOPOLOGY: Unknown
 - 1 ATGGGCGCG CATGAGAGAA GCCCAAACCA ATAACTACCC
- 10 AAAATGGAGA AAGTTCACGT
 - 61 TGACATCGAG GAAGATAGTC CCTTCCTCAG AGCATTACAA CGGAGCTTCC CGCAGTTTGA
 - 121 GGTAGAAGCC AAGCAGGTCA CAGATAATGA CCATGCTAAC GCCAGAGCGT TTTCGCATTT
- 15 181 GGCATCGAAA TTGATCGAGA CGGAGGTGGA ACCATCCGAT ACGATCCTAG ACATTGGAAG
 - 241 TGCGCCTGCC CGCAGAATGT ATTCCAAGCA TAAGTACCAT TGCATCTGTC CGATGAAATG
 - 301 TGCAGAAGAT CCGGACAGAC TGTTTAAGTA TGCAGCCAAG
 CTGAAGAAGA ACTGTAAAGA
 - 361 GATTACAGAT AAGGAACTGG ACAAGAAGAT GAAGGAGCTT GCGGAAGTCA TGAGCGACCC
 - 421 TGATCTCGAA ACTGAAACGA TTTGCCTTCA CGACGATGAA ACCTGTCGAT TTGAGGGTCA
- 25 481 AGTCGCAGTG TATCAGGATG TGTACGCGGT TGACGGACCG ACGAGCCTTT ACCATCAGGC
 - 541 CAACAAAGGG GTCAGAGTCG CCTATTGGAT AGGATTCGAC ACTACCCCTT TTATGTTTAA
- 601 GAACCTGGCT GGAGCATATC CCTCCTATTC GACCAACTGG
 30 GCCGACGAGA CCGTGTTAAC
 - 661 GGCTCGTAAT ATAGGCTTGT GCAGCTCCGA TGTGATGGAG CGGTCACGTA GAGGGATGTC
 - 721 CATCCTCAGA AAGAAATTTT TAAAACCATC CAATAACGTC TTGTTCTCTG TAGGATCTAC
- 781 CATCTACCAC GAGAAGCGAG ACTTACTAAG GAGTTGGCAC
 CTACCGTCCG TTTTTCACCT

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ACCCCCAGGG GGCTCATCAA

GCGGTGCTCT CCCCACAAGC

- 841 ACGTGGTAAG CAGAATTACA CTTGTCGGTG TGAGACTATA GTTAGTTGCG ACGGGTACGT 901 CGTCAAAAGG ATAGCTATTA GTCCAGGTCT GTACGGGAAA CCGTCGGGCT ACGCTGCCAC 961 GATGCATCGC GAGGGATTCT TGTGCTGCAA GGTGACGGAC ACGCTTGACG GGGAGAGGGT 1021 CTCTTTCCCC GTATGCACGT ACGTGCCAGC CACATTGTGC GACCAAATGA CAGGCATTCT 1081 GGCAACAGAT GTCAGTGCAG ACGACGCGCA GAAATTGCTG 10 GTTGGGCTCA ACCAGCGAAT 1141 AGTCGTAAAT GGCCGCACTC AAAGGAATAC TAACACAATG AAGAACTATT TATTACCCGT 1201 CGTGGCACAA GCATTTGCCA GATGGGCTAA AGAGTACACA GAAGATCAAG AAGATGAAAG 1261 ACCATTGGGG CTTAGGGACC GCCAGTTGGT AATGGGGTGT TGTTGGGCGT TCAGGAAACA 1321 CAAGATAACA TCAGTGTACA AACGACCAGA CACCCAAACG ATCATCAAGG TAAACAGCGA 1381 TTTCCACTCT TTCGTGCTGC CCAGAATTGG AAGCAACACC 20 TTGGAGATTG GGCTGAGGAC 1441 CAGGATCAGA AAACTACTGG AGGAACCTGT GGACAGACCA CCACTGATTA CCGCCGACGA 1501 CATACAGGAA GCCAAGAACG CGGCGGATGA GGCTAAGGAA GTCAAGGAGG CCGAAGAGCT 1561 CAGGGCAGCA TTACCACCGC TGTCTGCCGA TGTAGAGGAA CCTGCACTGG AGGCGGACGT 1621 TGACTTAATG CTGCAGGAGG CGGGAGCAGG ATCTGTCGAA
 - 1741 AGTACTGCGA AGTGAAAAAC TGACGTGCAT CCACCCGCTT GCAGAGCAAG TAATTGTAAT

1681 AGTCACCAGT TATGCAGGAG AAGACAAAAT TGGCTCTTAT

1801 CACACACTCT GGAAGGAAGG GTAGATACGC AGTTGAGCCT TACCACGGAA AAGTGGTAGT

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20

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ACCTCAGAGC ATGTAAATGT

1861 ACCCGAAGGG CAAGCTATAC CTGTCCAAGA CTTCCAAGCA CTCAGTGAGA GCGCCACAAT 1921 TGTGTACAAC GAACGTGAGT TCGTGAATAG GTATTTACAC CACATTGCCA CGCACGGTGG 1981 AGCTCTGAAC ACAGACGAGG AGTATTACAG AGTGGTGAAA CCTAGTGAAC ATGAAGGTGA 2041 GTACTTATAT GACATTGACA AGAAACAATG TGTGAAGAAA GAACTGGTTA CGGGACTAGG 2101 TTTGACAGGA GAACTTGTCG ATCCCCCCTT CCATGAATTT GCATATGAAA GCTTGAGGAC 2161 ACGTCCGCCC GCACCTTATC AAGTACCAAC TATAGGTGTG TATGGCGTTC CCGGGTCCGG 2221 GAAGTCTGGA ATTATTAAGA GCGCAGTCAC AAAGAAGGAC TTGGTAGTGA GTGCTAAGAA 2281 GGAGAACTGC GCCGAGATAA TAAGGGACGT CAAGAAGATG AAAGGGCTAG ACGTCAATGC 2341 CCGGACGGTG GACTCAGTGC TGTTAAATGG ATGCAAGCAC CCGGTCGAGA CTTTGTACAT 2401 TGATGAAGCC TTTGCATGCC ACGCCGGCAC TCTCAGGGCC TTAATAGCCA TTATACGCCC 2461 AAAGAAAGCA GTGCTATGTG GTGACCCAAA GCAATGTGGC TTCTTCAACA TGATGTGCCT 2521 GAAAGTGCAC TTCAACCATG AAATATGCAC TCAAGTTTTC CACAAAAGCA TATCTCGCAG 2581 GTGTACCAAG TCAGTGACGT CGGTAGTGTC CACACTGTTT TATGACAAAA GAATGAGAAC 2641 TACTAACCCA AGGGATTCCA AAATCGAAAT TGACACAACA GGAAGTACTA AACCAAAAA 2701 GGATGACTTG ATTCTCACAT GTTTTAGGGG ATGGGTTAAG 30 CAACTGCAAA TAGACTACAA 2761 AGGAAATGAA ATAATGACCG CGGCTGCCTC ACAGGGATTG ACGCGGAAAG GTGTCTATGC

2821 AGTTAGGTAC AAAGTTAACG AGAACCCATT GTACGCACCC

- 2881 GCTGTTGACC CGGACGGAAG ACAAGATTGT GTGGAAGACT CTTGCAGGGG ACCCGTGGAT
- 2941 AAAGACCCTG ACCGCGAAGT ACCCCGGAGA TTTCACTGCA ACAATGGAAG AATGGCAGGC
- 5 3001 AGAACATGAT GCCATCATGA GACACATCCT GGAGAAACCG GATCCCACGG ATGTCTTCCA
 - 3061 AAATAAAGCT AATGTTTGCT GGGCAAAGGC ACTTGTACCT GTGCTTAAGA CAGCCGGGAT
- 3121 AGATTTGACC ACAGAGCAGT GGAACACAGT GGATTACTTC
 10 AAAGAGGATA AGGCCCACTC
 - 3181 AGCTGAGATT GTCCTGAATC AGCTGTGCGT GCGATTCTTC GGTCTAGACT TAGATTCTGG
 - 3241 TTTGTTTTCC GCCCCACAG TTCCACTCTC CATTAGGAAC AACCATTGGG ACAACTCACC
- 15 3301 GTCACCCAAC ATGTACGGGT TGAATCAAGA AGTGGTCAGG CAACTATCAC GCAGGTACCC
 - 3361 TCAATTACCA CGTGCGGTGA CTACTGGGAG AGCATACGAC ATGAACACCG GTACTTTGCG
 - 3421 CAATTATGAT CCGCGCATAA ATTTAGTACC GGTGAACCGT CGTCTACCAC ATGCTCTCGT
 - 3481 GACGCAACAT GCTGATCATC CTCCCAGTGA TTTTTCCGCC TTTGTCAGTA AGCTTAAAGG
 - 3541 CAGAACGGTC CTAGTAGTTG GTGAGAAGAT GAGTATTTCA GGTAAGACGG TAGACTGGTT
- 25 3601 ATCTGAAACA CCTGATTCTA CTTTTAGGGC GCGCCTAGAT
 CTAGGCATAC CCAATGAACT
 - 3661 ACCGAAGTAC GATATCGTCT TCGTAAATGT AAGAACACAG TACCGCTACC ACCACTACCA
- 3721 GCAGTGTGAG GACCACGCCA TTAAGTTGAG CATGTTGACC
 30 AAGAAGGCCT GCCTGCACCT
 - 3781 GAACCCCGGA GGAACCTGTG TGAGCATTGG TTACGGCTAT GCGGACCGGG CCAGTGAAAG
 - 3841 CATCATAGGT GCAGTTGCTC GGCAGTTCAA GTTCTCGAGG GTATGCAAAC CGAAGGTGTC

- 3901 TAAGGAGGAG ACCGAAGTGC TATTTGTCTT CATTGGGTTC GATCGTAAAA CGCGAACCCA
- 3961 TAACCCATAC AAGCTCTCCT CCACCCTGAC CAATATTTAC ACCGGCTCGA GGCTCCATGA
- 5 4021 AGCTGGCTGC GCACCTTCGT ATCATGTAGT GCGCGGGGAT ATAGCCACTG CCACGGAAGG
 - 4081 AGTAATCGTT AATGCTGCCA ACAGCAAGGG CCAGCCAGGC AGTGGAGTGT GCGGAGCTCT
- 4141 GTACCGGAAG TACCCCGAAA GCTTCGATTT ACAACCAATA
 10 GAAGTGGGGA AAGCTAGATT
 - 4201 GGTCAAAGGT AACTCAAAAC ATCTCATTCA TGCAGTGGGG CCGAATTTTA ACAAAGTGTC
 - 4261 TGAAGTGGAA GGTGACAAAC AGCTGGCAGA AGCGTATGAA TCTATAGCCA GGATTATTAA
- 4321 TGACAACAAT TATAGATCTG TGGCTATTCC GCTTCTGTCC ACTGGAATAT TTGCCGGAAA
 - 4381 CAAGGATAGG TTAATGCAAT CCTTAAACCA TCTGTTAACG GCATTGGACA CAACAGACGC
- 4441 AGATGTGGCC ATATACTGCA GAGACAAGAA ATGGGAAGTG
 20 ACGTTGAAAG AGGTCGTAGC
 - 4501 CAGGAGAGA GCGGTAGAGG AGATATGTAT CTCCGAAGAT TCCTCCGTAG CAGAGCCGGA
 - 4561 TGCAGAGCTG GTTAGAGTTC ACCCTAAGAG CTCTTTGGCT GGAAGGAAAG GTTACAGCAC
- 25 4621 TAGCGATGGG AAGACATTCT CATATCTTGA AGGAACCAAA
 TTTCATCAGG CGGCGAAGGA
 - 4681 CATGGCAGAA ATTAACGCTA TGTGGCCTGC CGCTACAGAG GCTAATGAGC AGGTGTGCTT
- 4741 ATACATTCTG GGTGAAAGTA TGAGCAGTAT AAGATCCAAA
 30 TGCCCCGTTG AGGAGTCAGA
 - 4801 GGCATCCACC CCACCAAGTA CATTGCCTTG CTTGTGCATC CACGCTATGA CCCCGGAACG
 - 4861 GGTTCAGCGT TTGAAAGCCT CCCGCCCCGA ACAAATTACA GTTTGTTCTT CCTTCCCATT

10

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AGAAAGAAGT TACAACTGAA

GTTGAAAACA TGAAGGCCAT

4921 GCCGAAGTAC AGAATAACAG GAGTGCAGAA GATTCAATGT TCGCATCCTA TACTTTTCTC 4981 TCCTAAAGTA CCTGAGTACA TACACCCTAG AAAGTACCTT GCAGACGCAG CTTCCGCAAA 5041 CAATGGGGCA GCCGAATCAA CATCGGTGGA CGTGCAGCCA CAGCTGGAAG AGAGTCCTGA 5101 GAACACGGAA CAACTGGTGG AGGAGGAAGA CAGTATAAGC GTGCTGTCTG AGGATACACC 5161 ACACCAAGTG CACCAAGTGG AGGCTGAAGT GCATCGCTTC AGCGCAAGTG CTCAATCTTC 5221 GTCCTGGTCC ATTCCACGTG CATCCGACTT TGATGTCGAG AGTCTTTCCG TGCTCGAATC 5281 CCTGGGTGCT AATGATACAA TCAGCATGGA GTCGTCCTCA AACGAAACAG CTCTTGCTTT 5341 GCGGACCATT TTTAAGACTC CACCCATTCC AAGGCCTCGA GTGCAGAGCA CATCCACAGA 5401 CGTGGTTAGT ATCTCAGCAC TCGAGTCTTG TGACAGCACC AGCGATGCGC GTAGCATAGA 5461 CTCGGATGAA ACCGATGTTT CCATCTTTGA CAAAAGGTTG GAGTTCCTGG CCAGACCTGT 5521 TCCCGCACCG CGAACCAAAT TTAGGACTCC ACCCGTCCCG AAACCGCGTG CGCGGAGGCC 5581 ATTACATCCT TTGTCTAGTA GATCAAGCTC GCGCTCTAGC CTGGCGTCTA ATCCACCAGG 5641 TGTTAACCGA GTGATCACTA GGGAAGAATT TGAGTCCTTC GTTGCCCAAC AGCAATGACG 5701 GTTCGACGCG GGCGCGTACA TTTTCTCCTC GGATACTGGT CAAGGACATT TGCAACAAAA 5761 ATCAGTAAGG CAGACAGTAT TGTCTGAAGT GGTGCTAGAG 30 AGGACTGAGT TAGAGATCTC 5821 GTACGCCCCG CGCCTCGACC TGAACAAAGA AGAAATACTG

5881 CCCTACGCAA GCTAACCGGA GTAGATATCA GTCACGGAGG

- 5941 AACAACTAAG AGAATCTTAC AGGGATTAGG TCACTACCTG
 AAATCTGAAG GCAAGGTGGA
 6001 GTGCTATCGT ACGTTGCACC CCGTACCTTT GTATTCAGCA
 AGTGTGAACA GAGCGTTCTC
- 5 6061 CAGTCCCAAA GTTGCTGTTG AAGCATGTAA CGTTGTTCTG AAGGAAAATT TTCCGACAGT
 - 6121 GGCGTCGTAC TGCATAATAC CTGAGTACGA CGCCTACTTG GACATGGTGG GCGGTGCATC
- 6181 ATGTTGCTTG GATACGGCGA GTTTTTGCCC TGCCAAGTTG
 10 CGTAGCTTTC CGAAGAAACA
 - 6241 CGCATACCTC GAGCCCACCA TTCGGTCTGC AGTCCCATCA GCAATTCAGA ACACGCTGCA
 - 6301 AAATGTACTC GCAGCTGCCA CAAAGAGAAA CTGCAATGTG ACTCAAATGA GGGAGCTGCC
- 15 6361 TGTACTGGAT TCTGCGGCCT TCAATGTAGA GTGTTTTAAA AAATACGCTT GCAATAATGA
 - 6421 GTATTGGGAG ACCTACAAGA AGAATCCTAT TAGATTGACC GAGGAAAATG TGGTCAACTA
- 6481 TATAACCAAG TTAAAAGGGC CGAAGGCGGC TGCCCTGTAT
 20 GCAAAGACTC ATAATTTAGA
 - 6541 CATGCTGCAA GACATACCCA TGGACAGGTT TATTATGGAT TTAAAAAGAG ATGTCAAGGT
 - 6601 AACTCCAGGA ACCAAGCATA CCGAAGAAAG GCCTAAGGTC CAAGTAATCC AGGCTGCAGA
- 25 6661 TCCATTGGCT ACAGCATACC TATGTGGGAT TCATAGAGAA
 TTGGTGCGCA GACTGAACGC
 - 6721 AGTTCTGTTG CCCAACATAC ACACATTATT TGACATGTCT GCTGAGGATT TCGACGCCAT
- 6781 AATTGCCGAG CACTTCCAAC CAGGCGATTG GGTGTTAGAG
 30 ACAGACATAG CGTCATTCGA
 - 6841 TAAAAGCGAA GATGACGCGA TGGCTCTGAC GGCACTGATG ATCCTGGAAG ACCTCGGGGT
 - 6901 GGACCCAGAG CTGTTGACCC TAATCGAAGC GGCATTTGGC GAAATATCCT CCATTCACTT

- 6961 ACCAACCAAA ACTAAATTTA GGTTTGGAGC CATGATGAAA TCAGGGATGT TCTTGACTCT
- 7021 GTTTGTCAAT ACTGTGATCA ATATGGTCAT AGCTAGCAGA GTTCTGCGTG AGAGACTGAC
- 5 7081 AAACTCCCCT TGCGCCGCGT TCATTGGCGA CGACAATATC GTGAAAGGGG TTAAGTCCGA
 - 7141 CAAACTCATG GCCGATAGGT GCGCTACATG GTTGAACATG GAAGTCAAAA TCATCGACGC
- 7201 AGTGGTTGGC GAGAAAGCTC CCTACTTCTG TGGTGGGTTT
 10 ATTTTATGTG ACTCTGTGAC
 - 7261 CGGAACTGCA TGCCGTGTAG CAGACCCTTT GAAGAGATTA
 TTTAAGCTTG GAAAACCACT
 - 7321 GGCTGTGGAT GATGAACATG ATGATGACAG GCGTCGAGCA CTACAGGAGG AATCTGCCCG
- 7381 GTGGAACCGG GTGGGAATTT TTTCCGAGCT GTGCAAAGCC
 GTCGAGTCGC GATATGAAAC
 - 7441 AGTGGGCACG GCTGTCATTA TCATGGCCAT GACTACGCTC GCCAGCAGTG TCGAGTCGTT
 - 7501 CAGTTGTCTA AGAGGGGCTT CTATATCCCT CTACGGCTAA
 CCTGAATGGA CTGCGACGTA
 - 7561 GTCAAGTCCG CCGAAATGTT TCCTTATCAA CCAATGTACC CAATGCAGCC CATGCCCTTC
 - 7621 CGCAACCCTT TTGCGACTCC CAGAAGACCA TGGTTTCCAA GGACCGACCC CTTTTTAGCG
- 7681 ATGCAGGTGC AAGAGCTGGC AAGGTCCATG GCCAACTTGA
 CGTTCAAGCA ACGGCGAGAT
 - 7741 GTGCCGCCG AGGGTCCACC GGCTAAGAAG AAGAAGAAGG ACAACTCGCA ACAAGGTGGT
- 7801 CGGAATCAGA ATGGAAAGAA AAAGAACAAG CTAGTAAAGA
 30 AAAAGAAGAA GACAGGGCCA
 - 7861 CCACCCCAA AAAATACTGG TGGCAAAAAG AAAGTCAATA GGAAGCCAGG GAAGAGACAA
 - 7921 CGAATGGTTA TGAAGTTGGA GTCAGACAAG ACATTCCCTA
 TCATGCTAGA TGGAAAAGTT

GAGAACTGTA CACACACCCA

7981 AATGGATATG CATGCGTGGT CGGTGGCAAG CTGTTTAGAC CACTGCATGT GGAGGGTAAG 8041 ATTGACAATG ACGTGTTGTC CTCCCTCAAG ACCAAAAAGG CATCTAAGTA TGATCTGGAG 5 8101 TATGCTGATG TGCCGCAGAG CATGCGCGCA GACACATTTA AATACACTCA TGAAAAACCC 8161 CAGGGCTATT ACAGCTGGCA CCATGGAGCA GTACAGTATG AAAATGGCAG ATTCACAGTG 8221 CCCAAAGGAG TCGGAGCCAA AGGAGATAGC GGTCGCCCCA 10 TACTTGACAA CCAAGGGCGT . 8281 GTGGTCGCTA TTGTGCTTGG CGGAGTGAAT GAAGGCTCCA GAACGGCACT GTCTGTCGTG 8341 ACGTGGAACG AAAAAGGGGT TACAGTCAAA TACACCCCCG AGAATAGCGA GCAGTGGTCC 15 8401 CTGGTGACCA CCATGTGCCT GCTAGCCAAT GTCACGTTCC CGTGCACCCA ACCACCCATC 8461 TGCTACGACC GTAAGCCAGC AGAGACTTTG TCCATGCTCA GTCATAACAT AGACAATCCT 8521 GGTTATGACG AGTTGCTCGA AGCAGTACTG AAATGTCCAG 20 GCAGAGGCAA GAGGTCCACG 8581 GAGGAGCTGT TTAAGGAGTA CAAGTTAACA CGCCCGTACA TGGCCAGGTG CATCAGGTGT 8641 GCGGTCGGAA GTTGCCACAG CCCCATAGCC ATAGAGGCGG TAAGGAGCGA AGGGCACGAT 8701 GGCTATGTAC GACTCCAGAC CTCATCTCAG TATGGATTAG ACCCATCAGG AAACTTGAAA 8761 GGCAGAACCA TGAGGTATGA TATGCATGGA ACCATAGAAG AGATACCGTT GCATCAGGTG 8821 TCTGTTCATA CCTCACGTCC TTGCCACATA ATAGATGGGC 30 ATGGATACTT TCTGCTTGCC 8881 AGGTGCCCTG CAGGAGACTC CATAACTATG GAATTTAAGA AAGAATCAGT CACCCATTCC

8941 TGCTCTGTGC CCTACGAAGT AAAGTTTAAT CCTGCGGGAA

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CCCGCTTGCT GAAATGTGTG

GCGCCGGCGC TTACGAGCAC

9001 CCAGAGCACG GAGCTGAACA ACCTTGTCAC GTGTACGCTC ACGACGCACA AAATAGGGGA 9061 GCTTACGTGG AAATGCACCT TCCAGGATCC GAAGTGGACA GTACTTTACT GTCCATGAGC 9121 GGTAGTTCTG TTCATGTGAC TCCACCTGCC GGGCAAAGCG TCCAAGTGGA ATGCGAATGT 9181 GGTGGCACCA AGATCTCTGA AACCATCAAT TCAGCTAAAC AATACAGTCA GTGCTCAAAG 9241 ACATCTCAGT GCAGGGCATA CCGTACACAG AATGACAAAT GGGTGTACAA TTCGGATAAA 9301 CTGCCTAAAG CATCGGGAGA AACTCTCAAA GGCAAATTGC ATGTGCCCTT CGTACTGACC 9361 GAAGCGAAAT GCACAGTACC ATTGGCTCCA GAACCCATTA TCACCTTTGG GTTCCGCTCT 9421 GTGTCTCTGA AACTTCATCC TAAGAACCCC ACCTTCCTAA CCACGAGGCA GCTGGATGGA 9481 GAACCAGCTT ACACCCACGA ACTTATAACC CACCCTGTGG TGAGAAATTT CTCGGTTACA 9541 GAGAAAGGTT GGGAATTTGT GTGGGGAAAC CATCCGCCTC AAAGGTACTG GTCTCAAGAA 9601 ACTGCACCAG GTAATCCACA CGGACTACCA CACGAGGTGA TCACGCATTA CTATCACAGA 9661 TATCCCATGT CCACCATCCT CGGCTTATCA ATCTGTGCGG CGATAGTGAC GACATCCATT 9721 GCGGCATCCG TATGGCTGTT TTGCAAATCA CGGATTTCAT GCCTGACCCC CTATCGCTTG 9781 ACTCCGAATG CCAGCATGCC TCTGTGCTTA GCCGTCTTGT GCTGCGCACG CACAGCCAAA 9841 GCCGAAACTA CTTGGGAATC CCTAGATCAC CTCTGGAACC ACAACCAGCA GATGTTCTGG 9901 AGTCAGCTGC TAATCCCGCT AGCAGCACTG ATAGTTGCTA

9961 TGTTGCGTTG TGCCTTTTTT AGTCGTGGCC GGCGCCGTAG

- 10021 GCGACTACGA TGCCGAACCA AGTGGGGATC CCGTATAATA CCATTGTCAA CAGAGCGGGT
- 10081 TATGCACCTC TACCTATTAG CATAGTACCT ACTAAAGTGA AGCTGATTCC AACAGTGAAT
- 5 10141 CTTGAGTACA TTACATGCCA TTACAAGACT GGAATGGATT CACCCGCCAT TAAATGCTGC
 - 10201 GGCACTCAGG AGTGTTCTCC AACTTACAGG CCGGACGAGC AATGCAAAGT CTTCTCTGGA
- 10261 GTATACCCAT TTATGTGGGG AGGGGCGTAT TGCTTTTGCG
 10 ATACGGAGAA TACCCAGATA
 - 10321 AGCAAGGCGT ACGTGACGAA ATCGGAAGAT TGCGTCACCG ATCACGCCCA GGCATACAAA
 - 10381 GCACATACAG CCTCAGTCCA AGCCTTCTTA AATATTACAG TTGGAGGACA CTCAACGACA
- 15 10441 GCAGTGGTGT ATGTGAATGG AGAGACTCCC GTTAATTTTA
 ATGGAGTGAA GCTGACCGCG
 - 10501 GGCCCTCTGT CCACAGCCTG GTCGCCGTTC GACAAGAAGA TCGTGCAGTA CGCCGGGGAA
- 10561 ATTTATAACT ATGACTTTCC GGAATATGGA GCCGGCCACG
 20 CAGGAGCGTT TGGTGACATC
 - 10621 CAGGCTAGGA CGGTATCTAG TTCCGATGTA TACGCCAACA CAAACCTTGT GCTGCAGAGA
 - 10681 CCCAAAGCCG GAGCGATCCA TGTCCCGTAC ACCCAGGCCC CATCTGGGTA TGAACAATGG
- 25 10741 AAGAAAGATA AACCACCATC CCTCAAGTTC ACAGCCCCGT TCGGTTGTGA AATTTACACC
 - 10801 AACCCTATCC GTGCTGAAAA CTGCGCTGTG GGATCAATTC
- 10861 GATGCTCTGT TTACCAGGGT GTCCGAAACA CCGACATTAT
 30 CTGCTGCCGA GTGCACTCTG
 - 10921 AACGAGTGTG TATATTCATC CGACTTTGGC GGGATCGCTA CAGTCAAATA CTCGGCGAGC
 - 10981 AAGTCAGGCA AATGTGCAGT TCATGTACCC TCAGGCACGG CTACATTGAA AGAAGCCGCA

- 11041 GTCGAGTTGG CCGAACAGGG TTCGGCTACT ATACATTTTT CGACTGCCAG CATTCATCCG
- 11101 GAGTTTAGAC TCCAGATATG CACGTCTTAC GTTACGTGCA AAGGGGATTG TCACCCTCCG
- 5 11161 AAAGATCACA TTGTGACGCA TCCCCAATAC CACGCCCAGT CATTTACAGC TGCGGTATCA
 - 11221 AAAACCGCTT GGACGTGGTT AACATCCTTA CTGGGAGGGT CAGCTATAAT TATAATAATT
- 11281 GGACTTGTGT TAGCCACAGT TGTGGCTATG TATGTGCTGA 10 CCAACCAGAA ACATAATTAG
 - 11341 TATTAGCAGC GATTGGCATG CTGCTTGTAA AGTTTTATTA CAAATAACGT GCGGCAATTG
 - 11401 GCGAGCCGCT TTAATTAGAA TTTTATTTTC TTTTACCATA ATTGGATTTT GTTTTTAATA
- 15 11461 TTTC

(4) INFORMATION FOR SEQ ID NO:3:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 4003 base pairs
 - (B) TYPE: Nucleic acid
 - (C) STRANDEDNESS:Double (D) TOPOLOGY: Unknown
- Nucleotide Sequence of the BeAn8 26S Subgenomic 25 cDNA containing last portion of NS4, the 26S promoter, the complete BeAn8 structural genes and a portion of the 3' noncoding region.
- 30 1 CCCAGTAACT CTCTACGGCT GACCTGAATG GACTGTAACG TAGTTCAGTC
- 51 CGCAACCATG TTCCCTTACC AATCACCAAT GTTTCCAATG 35 CAACCAGCGC
 - 101 CTTTTCGCAA CCCGTACGCT CCTCCTAGAA GACCGTGGTT CCCTAGAACC
- 151 GATCCCTTCT TAGCCATGCA GGTGCAGGAG TTGGCCCGAT 40 CAATGGCGAG
 - 201 CTTGACGTTT AAACAGCGTC GAGATACGCC ACCCGAGGGG CCACCTGCTA

	GCAGGTTAAG	moco imoor	GCC1 CAACAGCA	AGG TAGCTCAGGC
5	301 AAAAAC CACCACCCCC	GAAACG GAAAAC	CGAA GAAGAAGA	AAA AGTAACGGAG
	351 AAAAAA CCTGGAAAAA	ATCAG AAGAGO	ACCA AGAAGAAG	GAC CAATAAGAAA
10	401 GACAAC TCCTATTTTG	CGGAT GGTTAT	GAAG TTAGAATC	AG ACAAAACATT
15	451 CTGGAT GAAAGCTATT	GGAA AAATTA	ATGG A'L'ATGCT'I	GC GTTGTTGGAG
	501 TCGACC	CATG CACGTG	GAAG GCAAAATT	GA CAATGAAACT
20	551 TGAAGA CGATGTTCCT	.CGAA GAAGGC	ATCC AAATACGA	CC TAGAGTATGC
25	601 CAAAGO AGCCTCAAGG	ATGC GAGCAG	ATAC CTTTAAAT	AC ACACATGATA
	651 GTATTA GGGAGATTCA	TAAT TGGCAT	CACG GCGCCGTG	CA GTATGAAAAT
30	701 CGGTGC CCCCATACTA	CGAA AGGTGT	GGGA GCGAAAGG.	AG ACAGTGGACG
	751 GATAAT TGAATGAAGG	CAAG GCAGAG	ICGT GGCCATTG	TG CTGGGAGGGG
35	801 ATCCAA GGGGTCACCG	GACA GCTTTG	ICCG TAGTTATG	TG GAATGAAAA
	851 TAAAAT TACCACCATG	ATAC ACCAGA	AAAC TGTGAGCA	AT GGTCACTAGT
10	901 TGTCTT CAATTTGCTA	CTCG CCGATG	TTAC GTTCCCTT	GT TCCACTCCAC
15	951 CGACCG AACATTGACA	AGCA CCCGCA	GAAA CCCTGATG.	AT GCTTAGCAAG
	1001 ATCCTG	GCTA TGATGA	ATTG CTGGAAGC	AG TGCTGAAATG
50	1051 CAGAAG	AGAT CTACGG	AGGA GTTATTTA.	AG GAGTACAAAC

	CACAGCCCGA
5	1151 TCGCTATAGA AGCAGTAAGA AGCGACGGCC ATGACGGCTA CATCCGAATA
	1201 CAGACATCAT CACAGTACGG TTTAGACCCC TCCGGGAACG TTAAGAGCAG
10	1251 AGTTATGAGG TATAATCTGT ATGGCAAGAT CGTAGAAGTT CCATTACATC
15	1301 AGGTTTCATT ACACACATCT CGGCCTTGCC ACATTATTGA TGGTCACGGA
	1351 TATTTCCTCC TCGCACGCTG CCCAGAGGGC GACTCTATCA CCATGGAGTT
20	1401 TAAGAAGGAT TCCGTCACCC ATTCCTGTTC AGTGCCTTAT GAAGTGAAAT
	1451 TCACACCCGT GGGCAGAGAA TTATATAGCC ATCCCCCAGA ACACGGCACA
25	1501 GAACATCCGT GCCGTGTGTA TGCCCACGAC GCCCAGCAAA AAGATGCGTA
30	1551 TGTGGAGATG CACCTGCCCG GGTCCGAAGT TGACAGTTCC CTGCTCTCCA
	1601 TGAGCGGTAG TGCGGTCCGG GTAACACCAC CATCAGGGCA AAGTGTCCTT
35	1651 GTGGAGTGCA ACTGTGGCTC CGCTGTGTCG GAAACCATAA ACACTGCAAA
	1701 ATCATACAGC CAATGCACAA AAACATCACA ATGCCGCGCG TACCGTCTGC
40	1751 AGAATGATAA GTGGGTATAC AATTCAGACA AACTTCCAAA GGCATCGGGA
45	1801 GAAACGCTGA AAGGGAAACT GCATGTACCT TATCTCCTTT CCGAAGCGAA
	1851 GTGTACCGTA CCTTTAGCAC CCGAGCCAAT AGTAACCTTC GGCTTTCGAT
50	1901 TCGTATCTTT GAAATTGCAT CCACGGAATC CGACATATTT GACTACCCGC

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	CAGAGCCAAC GAGCTAATTT
5	2001 AACTAGAAAT TTTACAGTGA CTGAGCATGG ATGGGGATAC GTTTGGGGTA
	2051 ATCACCCGCC TCAGAGGTAC TGGGCACAGG AGACAGCTCC AGGCAACCCG
10	2101 CATGGGCTGC CGCACGAGGT GATTACTCAT TACTATAACA GGTACCCAAT
15	2151 GTCCACGATT TTCGGACTAT CGATTTGCGC CGCAGTGGTA ACCACCTCAA
13	2201 TAGCCGCATC CACCTGGCTG TTGTGCAAGT CGAGAGTATC TTGTTTGACT
20	2251 CCGTATCGAC TGACCCCGAA TGCTCAGTTA CCTGTGTGTC TAGCCTTCCT
	2301 GTGCTGCGCG AGGACAGCCC GTGCAGAGAC CACATGGGAA TCACTAGACC
25	2351 ATTTATGGAA CCATAATCAA CAGATGTTCT GGAGTCAACT GCTCATTCCC
30	2401 CTAGCCGCGC TCATTGTGGT GACCCGCTTG CTGAAGTGCA TGTGTTGCGT
	2451 CGTTCCTTTT TTAGTCCTAG CAGGCGCCGC AAGCGTCGGC GCCTACGAAC
35	2501 ACGCAACCAC GATGCCGAAT CAGGTGGGGA TCCCGTATAA TACAGTAGTC
	2551 AACCGCGCAG GTGACGCACC ATTGGCAATC AGCATTATTC CAACCAAGAT
40	2601 ACGGCTAATT CCTACTTTGA ATTTAGAATA TATTACATGC CACTATAAGA
45	2651 CAGGATTAGA TTCACCTTTC ATTAAATGCT GCGGAACGCA GGAATGCCCC
	2701 CAAGTGAATA GACCCGATGA ACAGTGTAAA GTCTTCACGG GGGTGTATCC
50	2751 GTTTATGTGG GGAGGCGCCT ACTGCTTCTG TGACTCTGAA AACACGCAAA

2801 TTAGTCGAGC GTATGTGATG AAATCAGATG ACTGCTCAGC TGACCACGCC 2851 TTGGCCTACA AAGCTCATAC TGCCTCAGTC CAAGCTTTTC TGAATATAAC 2901 TGTGGGAGAG CAATCGACGA CAGCGGTAGT GTACGTGAAT GGAGAAACAC 10 2951 CGGTCAATTT TAACGGCATT AAATTGGTTG CAGGCCCTTT ATCAACTGCC 3001 TGGACCCCAT TTGATCGGAA AGTGGTGCAG TACGCCGGAG AGATCTACAA 15 3051 TTATGACTTC CCGGAGTACG GAGCTGGGCA TGCAGGGGCG TTCGGGGATC 3101 TTCAAGCCAG AACAATCACC AGTAATGACC TGTACGCCAA 20 CACGAATTTA 3151 GTATTGCAAA GACCCAACAC AGGCACCATC CATGTTCCTT ACACGCAGGC 25 3201 ACCGTCAGGC TTTGAGCAGT GGAAGAAAGA CAAACCACCA TCATTAAAGT 3251 ACACCGCACC ATTTGGGTGC GAAATTCATG TGAATCCCGT CAGAGCTGAG 30 3301 AATTGCGCAG TAGGATTTAT ACCATTAGCC TTCGACATAC CCGATGCCTT 3351 GTTTACCAGG GTGTCAGAAA CACCGACGTT GTCGAGCGCT 35 GAGTGCTCCT 3401 TGAATGAGTG TACATACTCA ACGGACTTTG GCGGGATCGC CACTGTCAAG 40 3451 TACTCGGCTA GTAAATCAGG CAAATGCGCA GTACATGTTC CCTCAGGCAC 3501 TGCAACTCTG AAAGAGTCAT TGGTGGAAGT GGTCGAACAA GGGTCAATGA 45 3551 CCCTTCACTT TTCGACCGCC AGTATACACC CAGAGTTTAA ATTGCAGATC 3601 TGTACGAAGG TACTCACATG TAAAGGCGAC TGTCATCCGC CTAAAGACCA 50

	3651 GCTGCGGTA	TATTGTGACG T	CACCCCCAGC	ACCACGCCCA	GACATTTACA
5	3701 GTCAGCAGT	CCAAGACCGC A	TTGGACGTGG	TTAACGTCAC	TCTTAGGAGG
	3751 TGTATGTGC	ATTATTATAA T	TTGGCCTTGT	ATTAGCAACT	GTTGTTGCTA
10	3801 AAGCTGCCT	GACCAACCAG A	AAACATAATT	AGACCACAGC	AGCGATTGGA
	3851 GTTTAGCAG	TTAGAAACAT C	GTAGCGGCAA	TTGGCAAGCC	GCCTATAAAT
15	3901 GCACGCCGC	AATTGGCAAG T		AAATTACCTA	GCGGCAATTG
20	3951 TTTTTAATA		TTATTTTCTT	TTACCAATAA	TTGGATTTTG
	4001	TTC			

What is claimed is:

- 1. A DNA comprising an isolated and purified western equine encephalitis (WEE) virus cDNA fragment coding for infectious western equine Encephalitis virus genome.
- 2. A DNA according to claim 1 wherein said WEE cDNA fragment is operably linked to a promoter such that said cDNA is transcribed.
 - 3. The DNA according to claim 1, wherein said cDNA fragment contains a deletion in the E3-E2 cleavage site.

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4. The DNA according to claim 3, wherein said cDNA fragment deletion is 5 amino acids at the E2-E2 cleavage site said amino acids being Arg Pro Lys Arg.

- 5. The DNA according to claim 4 wherein said cDNA fragment further contains a suppressor mutation.
- 6. The DNA according to claim 5 wherein said suppressor mutation is a substitution of glutamic acid at codon 182 of E2 to lysine.
- 7. The DNA according to claim 5 wherein said suppressor mutation is a substitution of glutamic acid at codon 181 of E2 to lysine.
 - 8. The DNA according to claim 2 wherein said promoter is T7 promoter of pBluescript KS and said DNA is pWE2000.

- 9. An infectious WEE virus RNA transcript encoded by the cDNA fragment of claim 1.
- 5 10. A DNA according to claim 5 wherein said WEE cDNA fragment is operably linked to a promoter such that said cDNA is transcribed.
- 11. An attenuated WEE virus RNA transcript
 10 encoded by the cDNA fragment of claim 5.
 - 12. A DNA according to claim 6 wherein said WEE cDNA fragment is operably linked to a promoter such that said cDNA is transcribed.

- 13. An attenuated WEE virus RNA transcript encoded by the cDNA fragment of claim 6.
- 14. A DNA according to claim 7 wherein said 20 WEE cDNA fragment is operably linked to a promoter such that said cDNA is transcribed.
 - $$15.\ An}$ attenuated WEE virus RNA transcript encoded by the cDNA fragment of claim 7.

- 16. Infectious WEE virus particles containing an RNA transcript according to claim 9.
- 17. Attenuated WEE virus particles
 30 containing an RNA transcript according to claim 11.
 - 18. Attenuated WEE virus particles containing an RNA transcript according to claim 13.

- 19. Attenuated WEE virus particles containing an RNA transcript according to claim 15.
- 20. A live attenuated western equine encephalitis (WEE) virus vaccine comprising attenuated western equine encephalitis virus according to claim 17.
- 21. A live attenuated western equine
 10 encephalitis (WEE) virus vaccine comprising attenuated
 Western Equine Encephalitis virus according to claim
 18.
- 22. A live attenuated western equine
 encephalitis (WEE) virus vaccine comprising attenuated
 western equine encephalitis virus according to claim
 19.
- 23. A DNA comprising an isolated and
 purified venezuelan equine encephalitis virus IE
 variant (VEE IE) cDNA fragment coding for infectious
 venezuelan equine encephalitis virus IE variant virus
 genome.
- 24. A DNA according to claim 23, wherein said cDNA fragment has the sequence of SEQ ID NO:2 or a portion thereof, or an allelic portion thereof.
- 25. The DNA according to claim 23, wherein 30 said cDNA fragment contains a deletion in the E3-E2 cleavage site.
 - 26. A DNA according to claim 25, wherein said cDNA fragment contains a deletion according to

claim 25 wherein said deletion is four amino acids said amino acids being Arg Gly Lys Arg.

- 27. A DNA fragment according to claim 26
 5 wherein said VEE IE cDNA further contains a suppressor mutation.
- 28. A DNA according to claim 23 wherein said VEE IE cDNA fragment is operably linked to a promoter such that said cDNA is transcribed.
 - 29. An infectious VEE IE virus RNA transcript encoded by the cDNA fragment of claim 23.
- 30. A DNA according to claim 27 wherein said VEE IE cDNA fragment is operably linked to a promoter such that said cDNA is transcribed.
- 31. An attenuated VEE IE virus RNA 20 transcript encoded by the cDNA fragment of claim 27.
 - 32. Infectious VEE IE virus particles containing an RNA transcript according to claim 29.
- 25 33. Attenuated VEE IE virus particles containing an RNA transcript according to claim 31.
 - 34. A live attenuated venezuelan equine encephalitis virus IE variant (VEE IE) virus vaccine comprising attenuated venezuelan equine encephalitis virus IE variant virus according to claim 33.
 - 35. A bivalent alphavirus vaccine comprising live attenuated western equine encephalitis (WEE)

 5 virus comprising an attenuating mutation and live

attenuated venezuelan equine encephalitis virus IE variant (VEE IE) comprising an attenuating mutation.

- 36. A bivalent alphavirus vaccine according to claim 35 wherein said live attenuated western equine encephalitis virus is chosen from the group consisting of attenuated WEE comprising a deletion in the E3-E2 cleavage site and a substitution of glutamic acid at codon 182 of E2 to lysine, and attenuated WEE comprising a deletion in the E3-E2 cleavage site and a substitution of glutamic acid at codon 181 of E2 to lysine.
- at nucleotide 13 of the 5' noncoding region of the WEE genome, and a deletion of the WEE genome.
- 38. A pharmaceutical formulation comprising attenuated WEE virus particles according to claim 17 in an effective immunogenic amount in a pharmaceutically acceptable carrier.
- 39. A pharmaceutical formulation comprising attenuated WEE virus particles according to claim 18 in an effective immunogenic amount in a pharmaceutically acceptable carrier.

40. A pharmaceutical formulation comprising attenuated WEE virus particles according to claim 19 in an effective immunogenic amount in a pharmaceutically acceptable carrier.

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41. A pharmaceutical formulation comprising attenuated VEE IE virus particles according to claim 33 in an effective immunogenic amount in a pharmaceutically acceptable carrier

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42. An attenuated chimeric virus comprising nonstructural sequences from a first alphavirus and structural sequences from a second alphavirus resulting in attenuation of the second alphavirus.

- 43. An attenuated chimeric virus according to claim 42 wherein wherein said first alphavirus is western equine encephalitis virus and said second alphavirus is chosen from the group consisting of:

 20 Sindbis virus, Aura virus, Barmah Forest virus, Bebaru Cabassou virus, Chikungunya virus, Everglades virus, Fort Morgan virus, Getah virus, Highlands J virus, Kyzylagach virus, Mayaro virus, Middelburg virus, Mucambo virus, Ndumu virus, O'nyong-nyong virus,

 25 Pixuna virus, Ross River virus, Sagiyama virus, Semliki Forest virus, SAAR87 virus, Tonate virus, Una virus, venezuelan equine encephalitis virus, and Whataroa virus.
- 30
- 44. An attenuated second virus according to claim 43 wherein said second alphavirus is eastern equine encephalitis virus.
- 45. An attenuated chimeric second virus according to claim 42 wherein said first alphavirus is

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Venezuelan equine encephalitis virus variant IE and said second alphavirus is chosen from the group consisting of: venezuelan equine encephalitis virus, western eqine encephalitis virus, eastern equine 5 encephalitis virus, Sindbis virus, Aura virus, Barmah Forest virus, Bebaru Cabassou virus, Chikungunya virus, Everglades virus, Fort Morgan virus, Getah virus, Highlands J virus, Kyzylagach virus, Mayaro virus, Middelburg virus, Mucambo virus, Ndumu virus, O'nyong-nyong virus, Pixuna virus, Ross River virus, Sagiyama virus, Semliki Forest virus, SAAR87 virus. Tonate virus, Una virusand Whataroa virus.

46. An attenuated chimeric second virus according to claim 44 wherein said second virus is 15 venezuelan equine encephalitis virus variant IA.

- 47. An attenuated chimeric second virus according to claim 42 wherein said first alphavirus is 20 venezuelan equine encephalitis virus variant IA and said second alphavirus is chosen from the group consisting of: Venezuelan equine encephalitis virus, western eqine encephalitis virus, eastern equine encephalitis virus, Sindbis virus, Aura virus, Barmah 25 Forest virus, Bebaru Cabassou virus, Chikungunya virus, Everglades virus, Fort Morgan virus, Getah virus, Highlands J virus, Kyzylagach virus, Mayaro virus, Middelburg virus, Mucambo virus, Ndumu virus, O'nyong-nyong virus, Pixuna virus, Ross River virus, Sagiyama virus, Semliki Forest virus, SAAR87 virus, 30 Tonate virus, Una virus, and Whataroa virus.
- 48. An attenuated chimeric second virus according to claim 47 wherein said second virus is 35 venezuelan equine encephalitis virus variant IE.

49. An attenuated chimeric second virus according to claim 47 wherein said second virus is Venezuelan Equine Encephalitis virus variant IIIA.

- 50. An attenuated virus vaccine comprising chimeric virus according to claim 42 wherein said vaccine is directed against said second alphavirus.
- 10 51. An inactivated western equine encephalitis virus (WEE) vaccine comprising attenuated WEE according to claim 17 wherein said attenuated WEE is inactivated.
- 52. An inactivated western equine encephalitis virus (WEE) vaccine comprising attenuated WEE according to claim 18 wherein said attenuated WEE is inactivated.
- 53. An inactivated western equine encephalitis virus (WEE) vaccine comprising attenuated WEE according to claim 19 wherein said attenuated WEE is further inactivated.
- 54. An inactivated venezuelan equine encephalitis virus variant IE (VEE IE) vaccine comprising attenuated VEE IE according to claim 33 wherein said attenuated VEE IE is further inactivated.
- 55. An inactivated eastern equine encephalitis virus (EEE) vaccine comprising attenuated EEE according to claim 44 wherein said attenuated virus is further inactivated.

56. An inactivated venezuelan equine encephalitis virus variant IA (VEE/IA) vaccine comprising attenuated VEE/IA according to claim 46 wherein said attenuated virus is further inactivated.

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57. An inactivated venezuelan equine encephalitis virus variant IIIA (VEE/IA) vaccine comprising attenuated VEE IIIA according to claim 49 wherein said attenuated virus is further inactivated.

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58. An inactivated venezuelan equine encephalitis virus variant IE (VEE IE) vaccine comprising attenuated VEE IE according to claim 48 wherein said attenuated virus is further inactivated.

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59. A method for expressing a protein said method comprising cloning a gene encoding said protein into a an attenuated virus replicon, said replicon chosen from the group consisting essentially of attenuated WEE and attenuated VEE IE wherein transcription of said replicon yields RNA capable of infecting a cell in which said protein is to be expressed.

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- $60.\ \mbox{A}$ method for the diagnosis of western equine encephalitis virus (WEE) infection comprising the steps of :
- (i) contacting a sample from an individual suspected of having a WEE infection with all or a unique portion of WEE; and
- (ii) detecting the presence or absence of a WEE infection by detecting the presence or absence of a complex formed between WEE and antibodies specific therefor in the sample.

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- 61. A method for the diagnosis of venezuelan equine encephalitis virus variant IE (VEE IE) infection comprising the steps of :
- (i) contacting a sample from an individual suspected of having a VEE IE infection with all or a unique portion of VEE IE; and
- (ii) detecting the presence or absence of aVEE IE infection by detecting the presence or absenceof a complex formed between VEE IE and antibodiesspecific therefor in the sample.
 - 62. A method for the diagnosis of western equine encephalitis virus (WEE) from a sample using the polymerase chain reaction, said method comprising:
 - (i) extracting RNA from the sample;
 - (ii) reverse transcribing the RNA of (i) to DNA;
 - (iii) contacting said DNA with
 - (a) at least four nucleotide triphosphates,
 - (b) a primer that hybridizes to WEE DNA, and
 - (c) an enzyme with polynucleotide synthetic activity,

under conditions suitable for the hybridization and extension of said first primer by said enzyme,

- whereby a first DNA product is synthesized with said DNA as a template therefor, such that a duplex molecule is formed;
- (iv) denaturing said duplex to release said first DNA product from said DNA;
- (v) contacting said first DNA product with a reaction mixture comprising:
 - (a) at least four nucleotide triphosphates,
 - (b) a second primer that hybridizes to said first DNA, and

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and

(c) an enzyme with polynucleotide synthetic activity,

under conditions suitable for the hybridization and extension of said second primer by said enzyme, whereby a second DNA product is synthesized with said first DNA as a template therefor, such that a duplex molecule is formed;

- (vi) denaturing said second DNA product from said first DNA product;
- 10 (vii) repeating steps iii-vi for a sufficient number of times to achieve linear production of said first and second DNA products;
 - (viii) fractionating said first and second DNA products generated from said WEE DNA; and
- 15 (ix) detecting said fractionated products for the presence or absence of WEE in a sample.
 - 63. A method for the diagnosis of venezuelan equine encephalitis variant IE virus (VEE IE) from a sample using the polymerase chain reaction, said method comprising:
 - (i) extracting RNA from the sample;
 - (ii) reverse transcribing said RNA of (i) to DNA;
 - (iii) contacting said DNA with
 - (a) at least four nucleotide triphosphates,
 - (b) a primer that hybridizes to VEE IE DNA,
 - (c) an enzyme with polynucleotide synthetic activity,
- under conditions suitable for the hybridization and extension of said first primer by said enzyme, whereby a first DNA product is synthesized with said DNA as a template therefor, such that a duplex molecule is formed;

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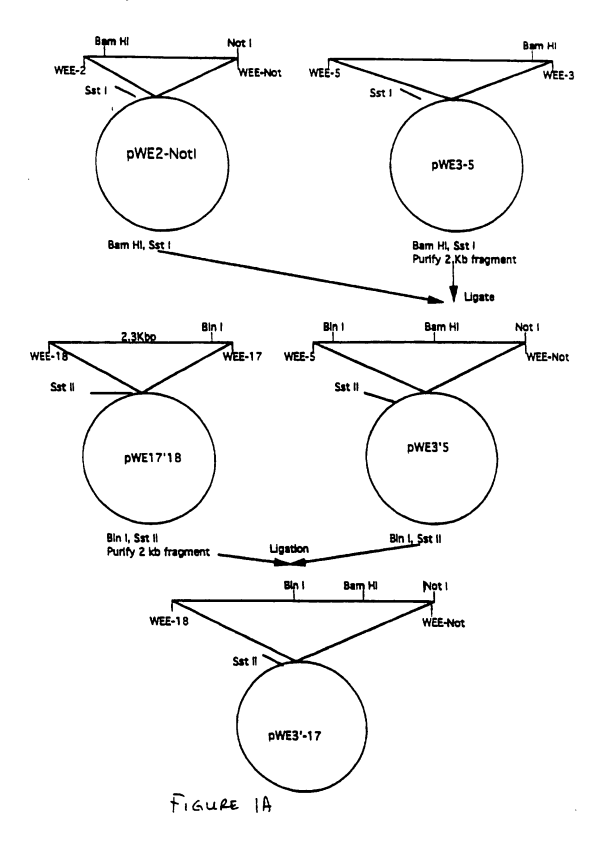
- (iv) denaturing said duplex to release said first DNA product from said DNA;
- (v) contacting said first DNA product with a reaction mixture comprising:
 - (a) at least four nucleotide triphosphates,
- (b) a second primer that hybridizes to said first DNA, and
- (c) an enzyme with polynucleotide synthetic activity,
- under conditions suitable for the hybridization and extension of said second primer by said enzyme, whereby a second DNA product is synthesized with said first DNA as a template therefor, such that a duplex molecule is formed;
- (vi) denaturing said second DNA product from said first DNA product;
 - (vii) repeating steps iii-vi for a sufficient number of times to achieve linear production of said first and second DNA products;
- 20 (viii) fractionating said first and second DNA products generated from said VEE IE DNA; and
 - (ix) detecting said fractionated products for the presence or absence of VEE IE in a sample.
- 25 64. A method for providing protective immunity against a second alphavirus to individuals with pre-existing immunity to a first alphavirus said method comprising administering to said individuals an effective amount of live attenuated second alphavirus.
 - 65. A method for providing protective immunity against a second alphavirus according to claim 64, wherein said live attenuated second alphavirus is attenuated western equine encephalitis virus.

- 66. The method of claim 65 wherein said western equine encephalitis (WEE) virus is chosen from the group consisting of: attenuated WEE having a substitution of glutamic acid at codon 181 of E2 to lysine, attenuated WEE having a substitution of glutamic acid at codon 181 of E2 to lysine. attenuated WEE having a C to T change at nucleotide 7 of the 5' noncoding region of the WEE genome, attenuated WEE having a A to G change at nucleotide 13 10 of the 5' noncoding region of the WEE genome, attenuated WEE having a T to A change at nucleotide 25 of the 5' noncoding region of the WEE genome, and attenuated WEE having a deletion of an A at nucleotide 22 of the 5' noncoding region of the WEE genome. 15
 - 67. A method for providing protective immunity against a second alphavirus according to claim 64, wherein said live attenuated second alphavirus is attenuated venezuelan equine encephalitis virus variant IE.
- 68. A WEE infection diagnostic kit comprising primers specific for WEE and ancillary reagents suitable for use in detecting the presence or absence of WEE in a mammalian sample.
- 69. A VEE IE infection diagnostic kit comprising primers specific for VEE IE and ancillary reagents suitable for use in detecting the presence or absence of VEE IE in a mammalian sample.
- 70. An isolated and purified Venezuelan equine encephalitis variant IIIA cDNA encoding the VEE 35 IIIA structural genes.

71. An amino acid fragment encoded by the cDNA fragment according to claim 70.

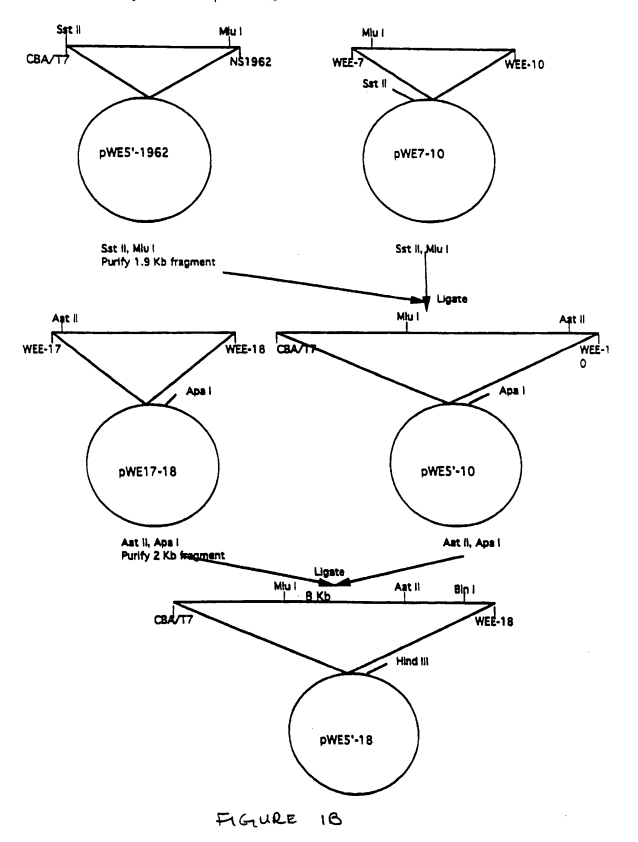
5 72. A polypeptide encoded by the amino acid fragment according to claim 71.

Asssembly 'm, _uie WE3'-17 1 / 7



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Assembly c noc 3 pWE5'-18 2 / 7



Assembly WELOOO full length cDNA clone of western equine encephalitis virus from modules pWES'-18 and pWE3'-17

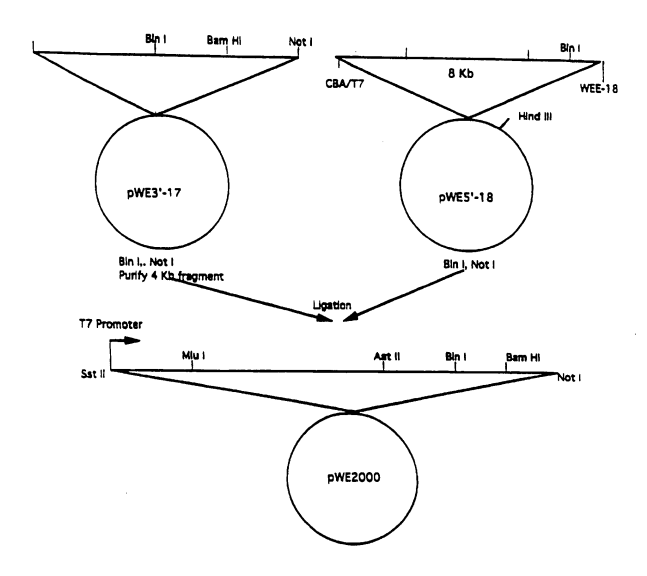


FIGURE 1C

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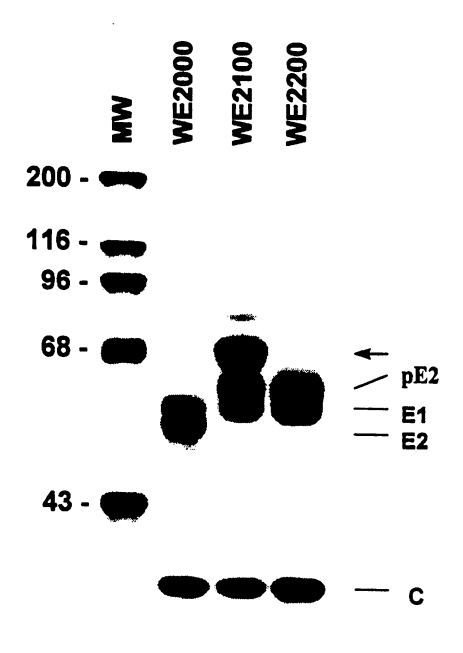


FIGURE 2

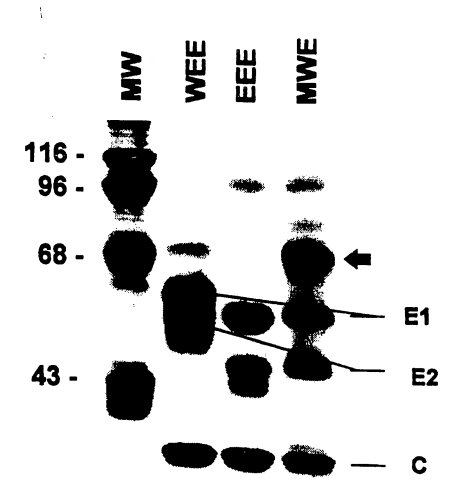
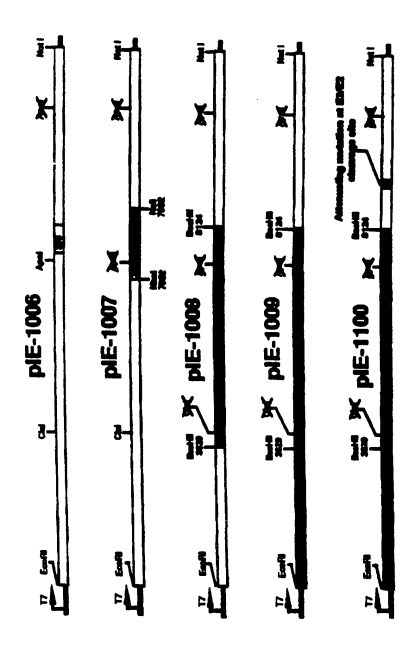


FIGURE 3



Floure 4

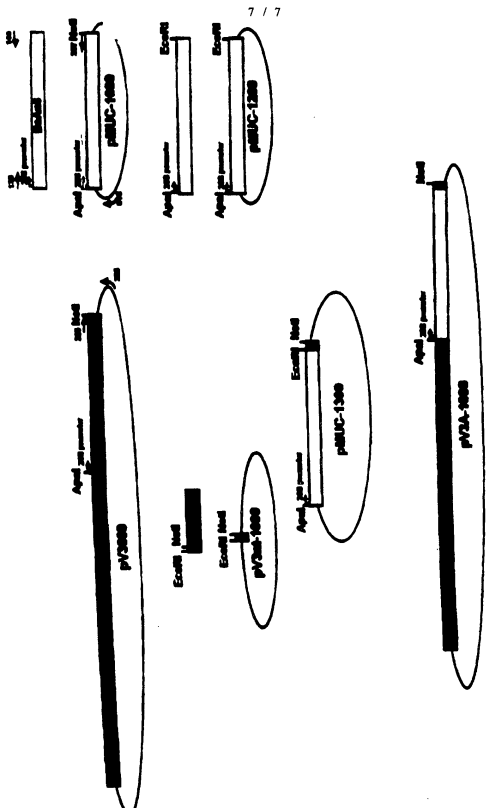


FIGURE 5

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	SSIFICATION OF SUBJECT MATTER			
IPC(6) US CL	:C12N 15/40, 7/01, 7/04, 15/86; A61K 39/193. :536/23.72; 435/235.1, 236, 69.1, 172.3; 424/205.1, 2	210.1		
According	to International Patent Classification (IPC) or to both	national classification and IPC		
	DS SEARCHED			
	ocumentation searched (classification system follower	ed by classification symbols)		
	536/23.72; 435/235.1, 236, 69.1, 172.3; 424/205.1, 2	•		
0.5.	330/23.72, 433/233.1, 230, 69.1, 172.3; 424/203.1, 2	18.1		
Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched				
	lata base consulted during the international search (n			
Aps, Bios sequenc?,	sis, Cab, Derwent WPI. Search terms: western, e dna, dnas, "E2", "E3", wee.	quine(w)enceph?, attenuat?, venezuelan	, genom?, ma. clon?,	
C. DOC	UMENTS CONSIDERED TO BE RELEVANT			
Category*	Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No.			
X	FRAIZER, G. et al. Isolation and preliminary characterization of mutants of Western equine encephalomyelitis virus with altered virulence in chickens. Biological Abstracts. 1985. Vol. 80, No. 5, page AB-506, Abstract no. 41179, see entire abstract.			
Y	US 5,185,440 A (DAVIS et al) 09 February 1993, see entire document, particularly column 7, line 66 through column 8, line 1.			
Y	US 5,505,947 A (JOHNSTON et al) 09 April 1996, see particularly column 3, lines 50-54, Abstract, and Examples 1 and 2.			
X Further documents are listed in the continuation of Box C. See patent family annex.				
* Special categories of cited documents: "T" later document published after the international filing date or priority				
"A" document defining the general state of the art which is not considered data and not in conflict with the application but cited to understand the principle or theory underlying the investigation.				
	E* extire document nublished on or after the international Clien data *X° document of particular relevance; the claimed invention cannot be			
"L" do	cument which may throw doubts on priority claim(s) or which is	considered novel or cannot be consider when the document is taken alone	red to involve an inventive step	
	cited to establish the publication date of another citation or other special reason (as specified) "Y" document of particular relevance; the claimed invention cannot be			
	considered to involve an inventive step when the document is			
	"P" document published prior to the international filing date but later than "&" document member of the same patent family			
	actual completion of the international search	Date of mailing of the international sea	rch report	
27 AUGU	27 AUGUST 1998 06 OCT 1998			
Name and mailing address of the ISA/US Commissioner of Patents and Trademarks Box PCT Westington D.C. 20221 MARY E. MOSHER, Ph.D.			Free	
Facsimile N	n, D.C. 20231 lo. (703) 305-3230	Telephone No. (703) 308-0196	H	

International application No. PCT/US98/10645

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C (Continua	tion). DOCUMENTS CONSIDERED TO BE RELEVANT		
Category*	Citation of document, with indication, where appropriate, of the relevant passages Relevant to		Relevant to claim No
Y	of Eastern and Western Equine Encephalomyelitis Viruses with Those of Other Alphaviruses and Related RNA Viruses. Virology. 1993. Vol. 197, pages 375-390. See figures 3 and 4.		1-3, 8, 9, 16, 59
?			1-3, 8, 9, 16, 59
\	STRAUSS, J.H. et al. The Alphaviruses: Gene Expressi Replication, and Evolution. Microbiological Reviews. S 1994. Vol. 58, No. 3, pages 491-562.	on, eptember	1-22, 37-40, 51- 53, 59
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International application No. PCT/US98/10645

Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)				
This international report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:				
Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:				
Claims Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:				
3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).				
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)				
This International Searching Authority found multiple inventions in this international application, as follows:				
Please See Extra Sheet.				
1. As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.				
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.				
As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:				
4. X No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.: 1-22, 37-40, 51-53, 59				
Remark on Protest The additional search fees were accompanied by the applicant's protest. No protest accompanied the payment of additional search fees.				
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International application No. PCT/US98/10645

BOX II. OBSERVATIONS WHERE UNITY OF INVENTION WAS LACKING This ISA found multiple inventions as follows:

This application contains the following inventions or groups of inventions which are not so linked as to form a single inventive concept under PCT Rule 13.1. In order for all inventions to be searched, the appropriate additional search fees must be paid.

Group 1, claims 1-22, 37-40, 51-53, 59, drawn to a first product DNA encoding an infectious genome of Western Equine Encephalitis (WEE) virus, RNA and virus encoded by DNA, vaccines and pharmaceuticals comprising virus encoded, and methods of use.

Group 2, claims 23-34, 41, 54, 59 drawn to second product DNA encoding an infectious genome of Venezuelan Equine Encephalitis IE variant (VEE IE) virus, RNA and virus encoded by DNA, vaccines and pharmaceuticals comprising virus encoded, and methods of use.

Group 3, claims 35-36, drawn to third product, combination vaccine comprising attenuated WEE and attenuated VEE IE

Group 4, claims 42-50, 55-58, drawn to fourth product, chimeric alphavirus and vaccine.

Group 5, claims 62, 68, drawn to fifth product, WEE specific primer kit, and method of use.

Group 6, claims 63, 69, drawn to sixth product, VEE IE specific primer kit, and method of use

Group 7, claims 70-72, drawn to seventh product, DNA encoding structural genes of VEE variant IIIA and products encoded.

Group 8, claims 60, drawn to method of diagnosing WEE using antigen.

Group 9, claims 61. drawn to method of diagnosing VEE IE using antigen.

Group 10, claims 64-67, drawn to method of vaccinating immune individual using a second alphavirus.

The inventions listed as Groups 1-10 do not relate to a single general inventive concept under PCT Rule 13.1 because, under PCT Rule 13.2, they lack the same or corresponding special technical features for the following reasons:

In groups 1 and 2, the special technical feature in each case is the DNA encoding an infectious genome of a particular virus; since groups 1 and 2 pertain to two distinct viruses, they involve different special technical features.

Group 3 does not have the corresponding special technical feature, because the attenuated viruses required in group 3 are not limited to those made by using the DNAs of groups 1 or 2; for example claim 35 encompasses combination vaccines using viruses attenuated by propagation in culture.

Group 4 does not have the corresponding special technical feature, because it is not limited to either of the viruses encoded by the DNAs of group 1 or group 2; for example claim 42 encompasses a chimera made from Sindbis and Semliki Forest viruses.

Groups 5 and 6 do not have the corresponding special technical feature, because they require small, specific primers, not infectious full-length genomic materials.

Group 7 does not have the corresponding special technical feature, because is drawn to a sub-genomic fragment of a third virus.

Groups 8 and 9 do not have the corresponding special technical feature, because they are not limited to materials of groups 1 or 2; for example claim 60 encompasses a method using viral antigens produced by propagation of native WEE in standard tissue culture.

Group 10 does not have the corresponding special technical feature because it is not limited to the viruses made using the DNAs of groups 1 or 2; for example claim 64 encompasses a method of vaccinating a Sindbis-immune monkey

INTERNATIONAL SEARCH REPORT International application No. PCT/US98/10645 against Semliki Forest virus.

Form PCT/ISA/210 (extra sheet)(July 1992)*